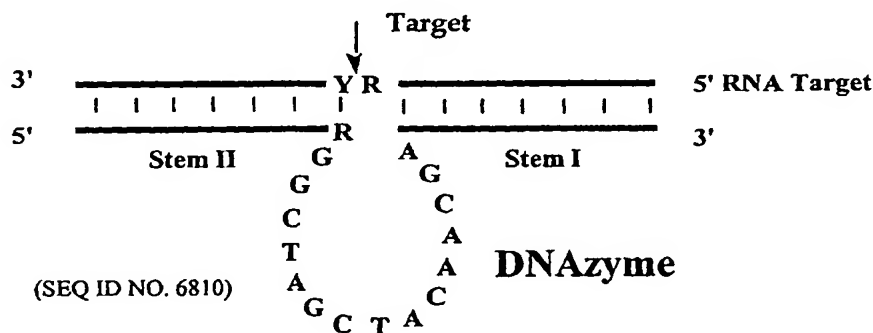




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DNAzyme Motif



(57) Abstract: The present invention relates to nucleic acid molecules, including enzymatic nucleic acid molecules, such as DNazymes (e.g. DNA enzymes, catalytic DNA), siRNA, aptamers, and antisense that modulate the expression of Ras genes such as K-Ras, H-Ras, and/or N-Ras. HIV genes such as HIV-1, and HER2 genes.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

DESCRIPTION

NUCLEIC ACID TREATMENT OF DISEASES OR CONDITIONS RELATED TO LEVELS OF RAS, HER2 AND HIV

This patent application claims priority from McSwiggen USSN 60/294,140, filed May 29, 2001, entitled "Enzymatic Nucleic Acid Treatment of Diseases or Conditions Related To Levels of HIV," McSwiggen USSN 60/296,249 filed June 6, 2001, entitled "Enzymatic Nucleic Acid Treatment of Diseases or Conditions Related to Levels of HER2," and McSwiggen USSN 60/318,471, filed September 10, 2001, entitled "Enzymatic Nucleic Acid Treatment of diseases or Conditions Related to Levels of RAS." Each of these applications is hereby incorporated by reference herein in its entirety including the drawings and tables.

Technical Field Of The Invention

The present invention relates to novel nucleic acid compounds and methods for the treatment or diagnosis of diseases or conditions related to levels of Ras gene expression, such as K-Ras, H-Ras, and/or N-Ras expression, HIV infection such as HIV-1, and HER2 gene expression.

Background Of The Invention

Transformation is a cumulative process whereby normal control of cell growth and differentiation is interrupted, usually through the accumulation of mutations affecting the expression of genes that regulate cell growth and differentiation.

The platelet derived growth factor (PDGF) system has served as a prototype for identification of substrates of the receptor tyrosine kinases. Certain enzymes become activated by the PDGF receptor kinase, including phospholipase C and phosphatidylinositol 3' kinase, Ras guanosine triphosphate (GTPase) activating protein (GAP) and src-like tyrosine kinases. GAP regulates the function of the Ras protein by stimulating the GTPase activity of the 21 kD Ras protein. Barbacid, 56 Ann. Rev. Biochem. 779, 1987. Microinjection of oncogenically activated Ras into NIH 3T3 cells has been shown to induce DNA synthesis. Mutations that cause oncogenic activation of Ras lead to accumulation of Ras bound to GTP, the active form of the molecule. These mutations block the ability of GAP to convert Ras to the inactive form. Mutations that impair the interactions of Ras with GAP also block the biological function of Ras.

While a number of Ras alleles exist (N-Ras, K-Ras, H-Ras) which have been implicated in carcinogenesis, the type most often associated with colon and pancreatic carcinomas is K-Ras. Enzymatic nucleic acid molecules which are targeted to certain regions of the K-Ras allelic mRNAs may also prove inhibitory to the function of the other allelic mRNAs of the N-Ras and H-Ras genes.

Scanlon, International PCT Publication Nos. WO 91/18625, WO 91/18624, and WO 91/18913 describes a ribozyme effective to cleave oncogene RNA from the H-Ras gene. This ribozyme is said to inhibit H-ras expression in response to exogenous stimuli. Reddy WO92/00080 describes the use of ribozymes as therapeutic agents for leukemias, such as chronic myelogenous leukemia (CML) by targeting specific portions of the BCR-ABL gene transcript.

Thompson *et al.*, International PCT publication No. WO 99/54459, describe nucleic acid molecules that modulate gene expression, including Ras gene expression.

Zhang *et al.*, 2000, *Gene Ther.*, 7, 2041; Takunaga *et al.*, 2000, *Br. J. Cancer.*, 83, 833; Zhang *et al.*, 2000, *Mol. Biotechnol.*, 15, 39; Irie *et al.*, 2000, *Mol. Urol.* 4, 61; Kijima and Scanlon, 2000, *Mol. Biotechnol.*, 14, 59; Funato *et al.*, 2000, *Cancer Gene Ther.*, 7, 495; Tsuchida *et al.*, 2000, *Cancer Gene Ther.*, 7, 373; Zhang *et al.*, 2000, *Methods Mol. Med.*, 35, 261; Irie *et al.*, 1999, *Antisense Nucleic Acid Drug Dev.*, 9, 341; Giannini *et al.*, 1999, *Nucleic Acids Res.*, 27, 2737; Fang *et al.*, 1999, *J. Med. Coll. PLA*, 14, 25; Tong *et al.*, 1998, *Methods Mol. Med.*, 11, 209; Ohkawa and Kashani-Sabet, 1998, *Methods Mol. Med.*, 11, 153; Scherr *et al.*, 1999, *Gene Ther.*, 6, 152; Tsuchida *et al.*, 1998, *Biochem. Biophys. Res. Commun.*, 252, 368; Scherr *et al.*, 1998, *Gene Ther.*, 5, 1227; Uhlmann *et al.*, European Patent Application EP 808898; Scherr *et al.*, 1997, *J. Biol. Chem.*, 272, 14304; Chang *et al.*, 1997, *J. Cancer Res. Clin. Oncol.*, 123, 91; Ohta *et al.*, 1996, *Nucleic Acids Res.*, 24, 938; Ohta *et al.*, 1994, *Ann. N.Y. Acad. Sci.*, 716, 242; and Funato *et al.*, 1994, *Biochem. Pharmacol.*, 48, 1471 all describe specific ribozymes targeting certain K-Ras, H-Ras, or N-Ras RNA sequences.

Todd, International PCT Publication Nos. WO 01/49877, WO 99/50452, and WO 99/45146 describes specific DNazymes targeting K-Ras for diagnostic applications.

Acquired immunodeficiency syndrome (AIDS) is thought to be caused by infection with the human immunodeficiency virus, for example HIV-1. Draper *et al.*, U.S. Patent Nos. 6,159,692, 5,972,704, 5,693,535, and International PCT Publication Nos. WO WO 93/23569,

WO 95/04818, describe enzymatic nucleic acid molecules targeting HIV. Todd *et al.*, International PCT Publication No. WO 99/50452, describe methods for using specific DNAzyme motifs for detecting the presence of certain HIV RNAs. Sriram and Banerjee, 2000, *Biochem J.*, 352, 667-673, describe specific RNA cleaving DNA enzymes targeting HIV-1. Zhang *et al.*, 1999, *FEBS Lett.*, 458, 151-156, describe specific RNA cleaving DNA enzymes used in the inhibition of HIV-1 infection.

HER2 (also known as neu, erbB2 and c-erbB2) is an oncogene that encodes a 185-kDa transmembrane tyrosine kinase receptor. HER2 is a member of the epidermal growth factor receptor (EGFR) family and shares partial homology with other family members. In normal adult tissues HER2 expression is low. However, HER2 is overexpressed in at least 25-30% of breast (McGuire, H.C. and Greene, M.I. (1989) The *neu* (c-erbB-2) oncogene. *Semin. Oncol.* 16: 148-155) and ovarian cancers (Berchuck, A. Kamel, A., Whitaker, R. *et al.* (1990)). Overexpression of her-2/neu is associated with poor survival in advanced epithelial ovarian cancer. *Cancer Research* 50: 4087-4091). Furthermore, overexpression of HER2 in malignant breast tumors has been correlated with increased metastasis, chemoresistance and poor survival rates (Slamon *et al.*, 1987 *Science* 235: 177-182). Because HER2 expression is high in aggressive human breast and ovarian cancers, but low in normal adult tissues, it is an attractive target for enzymatic nucleic acid-mediated therapy. McSwiggen *et al.*, International PCT Publication No. WO 01/16312 and Beigelman *et al.*, International PCT Publication No. WO 99/55857 describe enzymatic nucleic acid molecules targeting HER2. Thompson and Draper, US Patent No. 5,599,704, describes enzymatic nucleic acid molecules targeting HER2 (erbB2/neu) gene expression.

Summary Of The Invention

The present invention features nucleic acid molecules, including, for example, antisense oligonucleotides, siRNA, aptamers, decoys and enzymatic nucleic acid molecules such as DNAzyme enzymatic nucleic acid molecules, which modulate expression of nucleic acid molecules encoding Ras oncogenes, such as K-Ras, H-Ras, and N-Ras. In one embodiment, the invention features an enzymatic nucleic acid molecule comprising a sequence selected from the group consisting of SEQ ID NOs: 2329-4655.

In another embodiment, the invention features an enzymatic nucleic acid molecule comprising at least one binding arm having a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 1-2328.

5 In another embodiment, the invention features a siRNA molecule having complementarity to a sequence selected from the group consisting of SEQ ID NOs: 1-2328.

In another embodiment, the invention features an antisense molecule having complementarity to a sequence selected from the group consisting of SEQ ID NOs: 1-2328.

In another aspect of the invention, the nucleic acid of the invention is adapted to treat cancer.

10 In one embodiment, the enzymatic nucleic acid molecule of the invention has an endonuclease activity to cleave RNA having a K-Ras sequence.

In another embodiment, the enzymatic nucleic acid molecule of the invention has an endonuclease activity to cleave RNA having an H-Ras sequence.

15 In another embodiment, the enzymatic nucleic acid molecule of the invention has an endonuclease activity to cleave RNA having an N-Ras sequence.

In one embodiment, the siRNA molecule of the invention has RNA interference activity to K-Ras expression.

In another embodiment, the siRNA molecule of the invention has RNA interference activity to H-Ras expression.

20 In another embodiment, the siRNA molecule of the invention has RNA interference activity to N-Ras expression.

In one embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA is complementary to the RNA of K-Ras, H-Ras, and/or N-Ras gene. In another embodiment, a siRNA molecule of the invention comprises a double
25 stranded RNA wherein one strand of the RNA comprises a portion of a sequence of RNA of K-Ras, H-Ras, and/or N-Ras gene sequence. In yet another embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a non-nucleotide linker. Alternately, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a
30 nucleotide linker, such as a loop or stem loop structure.

In one embodiment, a single strand component of a siRNA molecule of the invention is from about 14 to about 50 nucleotides in length. In another embodiment, a single strand component of a siRNA molecule of the invention is about 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, or 28 nucleotides in length. In yet another embodiment, a single strand component of a siRNA molecule of the invention is about 23 nucleotides in length. In one embodiment, a siRNA molecule of the invention is from about 28 to about 56 nucleotides in length. In another embodiment, a siRNA molecule of the invention is about 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, or 52 nucleotides in length. In yet another embodiment, a siRNA molecule of the invention is about 46 nucleotides in length.

In one embodiment, the DNAzyme molecule of the invention is in a "10-23" configuration (see for example Santoro *et al.*, 1997, *PNAS*, 94, 4262 and Joyce *et al.*, US 5,807,718). In another embodiment, the DNAzyme comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 1-2328. In yet another embodiment, the DNAzyme comprises a sequence selected from the group consisting of SEQ ID NOs: 2329-4655.

In another embodiment, the nucleic acid molecule of the invention comprises between 12 and 100 bases complementary to a nucleic acid molecule having a K-Ras sequence. In yet another embodiment, the enzymatic nucleic acid comprises between 14 and 24 bases complementary to a nucleic acid molecule having a K-Ras sequence.

In another embodiment, the nucleic acid molecule of the invention comprises between 12 and 100 bases complementary to a nucleic acid molecule having an H-Ras sequence. In yet another embodiment, the nucleic acid molecule of the invention comprises between 14 and 24 bases complementary to a nucleic acid molecule having an H-Ras sequence.

In another embodiment, the nucleic acid molecule of the invention comprises between 12 and 100 bases complementary to a nucleic acid molecule having an N-Ras sequence. In yet another embodiment, the nucleic acid molecule of the invention comprises between 14 and 24 bases complementary to a nucleic acid molecule having an N-Ras sequence.

In yet another embodiment, the nucleic acid molecule of the invention is chemically synthesized. The nucleic acid molecule can comprise at least one 2'-sugar modification, at least one nucleic acid base modification, and/or at least one phosphate backbone modification.

In one embodiment, the invention features a mammalian cell comprising the nucleic acid molecule of the invention. In another embodiment, the mammalian cell of the invention is a human cell.

5 In another embodiment, the invention features a method of modulating K-Ras activity in a cell, comprising contacting the cell with the nucleic acid molecule of the invention, under conditions suitable for the modulation of K-Ras activity.

In another embodiment, the invention features a method of modulating H-Ras activity in a cell, comprising contacting the cell with the nucleic acid molecule of the invention, under conditions suitable for the modulation of H-Ras activity.

10 In another embodiment, the invention features a method of modulating N-Ras activity in a cell, comprising contacting the cell with the nucleic acid molecule of the invention, under conditions suitable for the modulation of N-Ras activity.

15 In another embodiment, the invention features a method of treatment of a subject having a condition associated with the level of K-Ras, comprising contacting cells of the subject with the nucleic acid molecule of the invention, under conditions suitable for the treatment.

20 In another embodiment, the invention features a method of treatment of a subject having a condition associated with the level of H-Ras, comprising contacting cells of the subject with the nucleic acid molecule of the invention, under conditions suitable for the treatment.

In another embodiment, the invention features a method of treatment of a subject having a condition associated with the level of N-Ras, comprising contacting cells of the subject with the nucleic acid molecule of the invention, under conditions suitable for the treatment.

25 In one embodiment, a method of treatment of the invention further comprises the use of one or more drug therapies under conditions suitable for the treatment.

30 In another embodiment, the invention features a method of cleaving RNA having a K-Ras sequence comprising contacting the K-Ras RNA with the enzymatic nucleic acid molecule of the invention under conditions suitable for the cleavage, for example, where the cleavage is carried out in the presence of a divalent cation, such as Mg²⁺.

In another embodiment, the invention features a method of cleaving RNA having a H-Ras sequence comprising contacting the H-Ras RNA with the enzymatic nucleic acid molecule of the invention under conditions suitable for the cleavage, for example, where the cleavage is carried out in the presence of a divalent cation, such as Mg^{2+} .

- 5 In another embodiment, the invention features a method of cleaving RNA having an N-Ras sequence comprising contacting the N-Ras RNA with the enzymatic nucleic acid molecule of the invention under conditions suitable for the cleavage, for example, where the cleavage is carried out in the presence of a divalent cation, such as Mg^{2+} .

10 In one embodiment, the nucleic acid molecule of the invention comprises a cap structure, for example, a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative, wherein the cap structure is at the 5'-end, 3'-end, or both the 5'-end and the 3'-end of the nucleic acid molecule.

15 In another embodiment, the invention features an expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of the invention in a manner that allows expression of the nucleic acid molecule. For example, the invention features an expression vector comprising a nucleic acid encoding a DNAzyme in a manner that allows expression of the DNAzyme.

In yet another embodiment, the invention features a mammalian cell, for example a human cell, comprising an expression vector of the invention.

20 In another embodiment, the expression vector of the invention further comprises a sequence for a nucleic acid molecule complementary to an RNA having K-Ras sequence.

In another embodiment, the expression vector of the invention further comprises a sequence for a nucleic acid molecule complementary to an RNA having H-Ras sequence.

25 In another embodiment, the expression vector of the invention further comprises a sequence for a nucleic acid molecule complementary to an RNA having N-Ras sequence.

30 In one embodiment, an expression vector of the invention comprises a nucleic acid sequence encoding two or more nucleic acid molecules of the invention, which can be the same or different. In another embodiment, an expression vector of the invention further comprises a sequence encoding an antisense nucleic acid molecule complementary to an RNA having a K-Ras, H-Ras or N-Ras sequence.

In another embodiment, the invention features a method for treating cancer, for example colorectal cancer, bladder cancer, lung cancer, pancreatic cancer, breast cancer, or prostate cancer, comprising administering to a subject a nucleic acid molecule of the invention under conditions suitable for the treatment. A method of treatment of cancer of the invention can further comprise administering to a patient one or more other therapies, for example, monoclonal antibody therapy, such as Herceptin (trastuzumab); chemotherapy, such as paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, Leucovorin, Irinotecan (CAMPTOSAR® or CPT-11 or Camptothecin-11 or Campto), Carboplatin, edatrexate, gemcitabine, or vinorelbine; radiation therapy, or analgesic therapy and/or any combination thereof.

In another embodiment, the invention features a composition comprising a nucleic acid molecule of the invention in a pharmaceutically acceptable carrier.

In one embodiment, the invention features a method of administering to a cell, for example a mammalian cell or human cell, the nucleic acid molecule of the invention comprising contacting the cell with the nucleic acid molecule under conditions suitable for administration. The method of administration can be in the presence of a delivery reagent, for example a lipid, cationic lipid, phospholipid, or liposome.

The present invention features an enzymatic nucleic acid molecule which modulates expression of a nucleic acid molecule encoding a human immunodeficiency virus (HIV), for example HIV-1, HIV-2, and related viruses such as FIV-1 and SIV-1, or a HIV gene, for example LTR, nef, vif, tat, or rev, wherein the enzymatic nucleic acid molecule comprises a DNAzyme configuration.

The invention also features an enzymatic nucleic acid molecule which modulates expression of a nucleic acid molecule encoding HIV or a component of HIV such as net, vif, tat, or rev, wherein the enzymatic nucleic acid molecule is in a Inozyme, G-cleaver, Zinzyme, DNAzyme or Amberzyme configuration.

The present invention also features a siRNA molecule which modulates expression of a nucleic acid molecule encoding a human immunodeficiency virus (HIV), for example HIV-1, HIV-2, and related viruses such as FIV-1 and SIV-1, or a HIV gene, for example LTR, nef, vif, tat, or rev.

The present invention features an enzymatic nucleic acid molecule comprising a sequence selected from the group consisting of SEQ ID NOs. 6727-6799. The invention also features an enzymatic nucleic acid molecule comprising at least one binding arm wherein one

or more of said binding arms comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6642-6726. In addition, the present invention features a siRNA nucleic acid molecule comprising sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 1-76 and 140-148.

5 In another embodiment, the siRNA molecule of the invention has RNA interference activity to HIV-1 expression and/or replication.

In one embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA is complementary to the RNA of HIV-1 genome or genes. In another embodiment, a siRNA molecule of the invention comprises a double
10 stranded RNA wherein one strand of the RNA comprises a portion of a sequence of HIV-1 genome or gene sequence. In yet another embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a non-nucleotide linker. Alternately, a siRNA molecule of the invention comprises a double
15 stranded RNA wherein both strands of RNA are connected by a nucleotide linker, such as a loop or stem loop structure.

In one embodiment, a single strand component of a siRNA molecule of the invention is from about 14 to about 50 nucleotides in length. In another embodiment, a single strand component of a siRNA molecule of the invention is about 14, 15, 16, 17, 18, 19, 20, 21, 22,
20 23, 24, 25, 26, 27, or 28 nucleotides in length. In yet another embodiment, a single strand component of a siRNA molecule of the invention is about 23 nucleotides in length. In one embodiment, a siRNA molecule of the invention is from about 28 to about 56 nucleotides in length. In another embodiment, a siRNA molecule of the invention is about 40, 41, 42, 43,
25 44, 45, 46, 47, 48, 49, 50, 51, or 52 nucleotides in length. In yet another embodiment, a siRNA molecule of the invention is about 46 nucleotides in length.

25 In one embodiment, a nucleic acid molecule of the invention is adapted to treat HIV infection or acquired immunodeficiency syndrome (AIDS).

In another embodiment, the enzymatic nucleic acid molecule of the invention has an endonuclease activity to cleave RNA having HIV sequence.

In yet another embodiment, the enzymatic nucleic acid molecule of the invention is in
30 an Inozyme, Zinzyme, G-cleaver, Amberzyme, DNazyme or Hammerhead configuration.

In another embodiment, the Inozyme of the invention comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6648-6655, or comprises a sequence selected from the group consisting of SEQ ID NOs. 6733-6740.

5 In another embodiment, the Zinzyme of the invention comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6663 and 6723-6726, or comprises a sequence selected from the group consisting of SEQ ID NOs 6741-6748 and 6795-6799.

10 In another embodiment, the Amberzyme of the invention comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6688, or comprises a sequence selected from the group consisting of SEQ ID NOs. 6762-6789.

In another embodiment, the DNAzyme of the invention comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6668 and 6718-6722, or comprises a sequence selected from the group consisting of SEQ ID NOs. 6749-6761 and 6790-6794.

15 In another embodiment, the Hammerhead of the invention comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6642-6647, or comprises a sequence selected from the group consisting of SEQ ID NOs 6727-6732.

In one embodiment, a nucleic acid molecule of the invention comprises between 12 and 100 bases complementary to a RNA sequence encoding HIV genome, RNA, and/or proteins.

20 In another embodiment, a nucleic acid molecule of the invention comprises between 14 and 24 bases complementary to a RNA sequence encoding HIV genome, RNA, and/or proteins.

In yet another embodiment, a nucleic acid molecule of the invention is chemically synthesized. A nucleic acid molecule of the invention can comprise at least one 2'-sugar modification, at least one nucleic acid base modification, and/or at least one phosphate backbone modification.

25 The present invention features a mammalian cell including a nucleic acid molecule of the invention. In one embodiment, the mammalian cell of the invention is a human cell.

The invention features a method of reducing HIV activity in a cell, comprising contacting the cell with a nucleic acid molecule of the invention, under conditions suitable for the reduction of HIV activity.

The invention also features a method of treating a subject having a condition associated with the level of HIV, comprising contacting cells of the subject with a nucleic acid molecule of the invention, under conditions suitable for the treatment.

5 In one embodiment, methods of treatment contemplated by the invention comprise the use of one or more drug therapies under conditions suitable for the treatment.

The invention features a method of cleaving RNA comprising a HIV nucleic acid sequence comprising contacting an enzymatic nucleic acid molecule of the invention with the RNA under conditions suitable for the cleavage. In one embodiment, the cleavage contemplated by the invention is carried out in the presence of a divalent cation, for example
10 Mg^{2+} .

The present invention features a method for treatment of acquired immunodeficiency syndrome (AIDS) or an AIDS related condition, for example Kaposi's sarcoma, lymphoma, cervical cancer, squamous cell carcinoma, cardiac myopathy, rheumatic disease, or opportunistic infection, comprising administering to a subject a nucleic acid molecule of the
15 invention under conditions suitable for the treatment.

In one embodiment, nucleic acid molecule of the invention comprises at least five ribose residues, at least ten 2'-O-methyl modifications, and a 3'- end modification, for example a 3'-3' inverted abasic moiety.

In another embodiment, a nucleic acid molecule of the invention further comprises
20 phosphorothioate linkages on at least three of the 5' terminal nucleotides.

In yet another embodiment, a DNAzyme of the invention comprises at least ten 2'-O-methyl modifications and a 3'-end modification, for example a 3'-3' inverted abasic moiety.

In a further embodiment, the DNAzyme of the invention further comprises phosphorothioate linkages on at least three of the 5' terminal nucleotides.

25 In another embodiment, other drug therapies of the invention comprise antiviral therapy, monoclonal antibody therapy, chemotherapy, radiation therapy, analgesic therapy, or anti-inflammatory therapy.

In yet another embodiment, antiviral therapy of the invention comprises treatment with AZT, ddC, ddI, d4T, 3TC, Ribavirin, delvaridine, nevirapine, efavirenz, ritonavir, saquinavir,
30 indinavir, amprenavir, nelfinavir, or lopinavir.

The invention features a composition comprising a nucleic acid molecule of the invention in a pharmaceutically acceptable carrier.

In one embodiment, the invention features a method of administering to a cell, for example a mammalian cell or human cell, an enzymatic nucleic acid molecule of the invention comprising contacting the cell with the enzymatic nucleic acid molecule under conditions suitable for the administration. The method of administration can be in the presence of a delivery reagent, for example a lipid, cationic lipid, phospholipid, or liposome.

The present invention features enzymatic nucleic acid molecules which modulate expression of nucleic acid molecules encoding HER2. The present invention also features siRNA molecules which modulate the expression of nucleic acid molecules encoding HER2.

In another embodiment, the invention features a siRNA molecule having complementarity to a sequence selected from the group consisting of SEQ ID NOs: 4656-5643 and 6632-6636.

In one embodiment, the invention features an enzymatic nucleic acid molecule comprising a sequence selected from the group consisting of SEQ ID NOs: 5644-6631 and 6637-6641.

In another embodiment, the invention features an enzymatic nucleic acid molecule comprising at least one binding arm having a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 4656-5643 and 6632-6636.

In yet another embodiment, a nucleic acid of the invention is adapted to treat cancer.

In another embodiment, an enzymatic nucleic acid molecule of the invention has an endonuclease activity to cleave RNA having HER2 sequence.

In another embodiment, the siRNA molecule of the invention has RNA interference activity to N-Ras gene expression.

In one embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA is complementary to the RNA of HER2 gene. In another embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA comprises a portion of a sequence of RNA having of HER2 gene sequence. In yet another embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a non-nucleotide linker. Alternately, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a nucleotide linker, such as a loop or stem loop structure.

In one embodiment, a single strand component of a siRNA molecule of the invention is from about 14 to about 50 nucleotides in length. In another embodiment, a single strand component of a siRNA molecule of the invention is about 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, or 28 nucleotides in length. In yet another embodiment, a single strand component of a siRNA molecule of the invention is about 23 nucleotides in length. In one embodiment, a siRNA molecule of the invention is from about 28 to about 56 nucleotides in length. In another embodiment, a siRNA molecule of the invention is about 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, or 52 nucleotides in length. In yet another embodiment, a siRNA molecule of the invention is about 46 nucleotides in length.

In one embodiment, a DNzyme molecule of the invention is in a "10-23" configuration. In another embodiment, a DNzyme of the invention comprises a sequence complementary to a sequence having SEQ ID NOs: 4656-5643 and 6632-6636. In yet another embodiment, a DNzyme molecule of the invention comprises a sequence having SEQ ID NOs: 5644-6631 and 6637-6641.

In another embodiment, a nucleic acid molecule of the invention comprises between 12 and 100 bases complementary to a nucleic acid molecule having HER2 sequence. In yet another embodiment, a nucleic acid molecule of the invention comprises between 14 and 24 bases complementary to a nucleic acid molecule having HER2 sequence.

In yet another embodiment, a nucleic acid molecule of the invention is chemically synthesized. A nucleic acid molecule of the invention can comprise at least one 2'-sugar modification, at least one nucleic acid base modification, and/or at least one phosphate backbone modification.

In one embodiment, the invention features a mammalian cell comprising a nucleic acid molecule of the invention. In another embodiment, the mammalian cell of the invention is a human cell.

In another embodiment, the invention features a method of reducing HER2 activity in a cell, comprising contacting the cell with the nucleic acid molecule of the invention, under conditions suitable for the reduction of HER2 activity.

In another embodiment, the invention features a method of treatment of a subject having a condition associated with the level of HER2, comprising contacting cells of the subject with the nucleic acid molecule of the invention, under conditions suitable for the treatment.

In one embodiment, a method of treatment of the invention further comprises the use of one or more drug therapies under conditions suitable for the treatment.

5 In another embodiment, the invention features a method of cleaving RNA having HER2 sequence comprising contacting an enzymatic nucleic acid molecule of the invention with the RNA under conditions suitable for the cleavage, for example, where the cleavage is carried out in the presence of a divalent cation, such as Mg^{2+} .

10 In one embodiment, a nucleic acid molecule of the invention comprises a cap structure, for example a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative, wherein the cap structure is at the 5'-end, 3'-end, or both the 5'-end and the 3'-end of the enzymatic nucleic acid molecule.

In another embodiment, the invention features an expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of the invention, for example a DNAzyme or siRNA molecule, in a manner that allows expression of the nucleic acid molecule.

15 In yet another embodiment, the invention features a mammalian cell, for example a human cell, comprising an expression vector of the invention.

In another embodiment, an expression vector of the invention further comprises a sequence for a nucleic acid molecule complementary to a nucleic acid molecule having HER2 sequence.

20 In one embodiment, an expression vector of the invention comprises a nucleic acid sequence encoding two or more nucleic acid molecules, which can be the same or different. In another embodiment, an expression vector of the invention further comprises a sequence encoding an antisense nucleic acid molecule complementary to a nucleic acid molecule having a HER2 sequence.

25 In another embodiment, the invention features a method for treating cancer, for example breast cancer or ovarian cancer, comprising administering to a subject a nucleic acid molecule of the invention under conditions suitable for the treatment. A method of treatment of cancer of the invention can further comprise administering to a patient one or more other therapies, for example, monoclonal antibody therapy, such as Herceptin (trastuzumab);
30 chemotherapy, such as paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, Leucovorin, Irinotecan (CAMPTOSAR® or CPT-11 or Camptothecin-11 or Campto), Carboplatin, edatrexate,

gemcitabine, or vinorelbine; radiation therapy, or analgesic therapy and/or any combination thereof.

In another embodiment, the invention features a composition comprising a nucleic acid molecule of the invention in a pharmaceutically acceptable carrier.

5 : In one embodiment, the invention features a method of administering to a cell, for example a mammalian cell or human cell, a nucleic acid molecule of the invention comprising contacting the cell with the nucleic acid molecule under conditions suitable for administration. The method of administration can be in the presence of a delivery reagent, for example a lipid, cationic lipid, phospholipid, or liposome.

10

Detailed Description of the Invention

First the drawings will be described briefly.

Drawings

Figure 1 shows examples of chemically stabilized ribozyme motifs. **HH Rz**, represents
15 hammerhead ribozyme motif (Usman *et al.*, 1996, *Curr. Op. Struct. Bio.*, 1, 527); **NCH Rz**
represents the NCH ribozyme motif (Ludwig *et al.*, International PCT Publication No. WO
98/58058 and US Patent Application Serial No. 08/878,640); **G-Cleaver**, represents G-
cleaver ribozyme motif (Kore *et al.*, 1998, *Nucleic Acids Research* 26, 4116-4120, Eckstein *et*
20 *al.*, US 6,127,173). **N** or **n**, represent independently a nucleotide which can be same or
different and have complementarity to each other; **rI**, represents ribo-Inosine nucleotide;
arrow indicates the site of cleavage within the target. Position 4 of the HH Rz and the NCH
Rz is shown as having 2'-C-allyl modification, but those skilled in the art will recognize that
this position can be modified with other modifications well known in the art, so long as such
modifications do not significantly inhibit the activity of the ribozyme.

25 **Figure 2** shows an example of the Amberzyme ribozyme motif that is chemically
stabilized (see for example Beigelman *et al.*, International PCT publication No. WO
99/55857 and US Patent Application Serial No. 09/476,387.).

Figure 3 shows an example of a Zinzyme A ribozyme motif that is chemically
stabilized (see for example Beigelman *et al.*, International PCT publication No. WO
30 99/55857 and US Patent Application Serial No. 09/918,728).

Figure 4 shows an example of a DNAzyme motif described by Santoro *et al.*, 1997, *PNAS*, 94, 4262 and Joyce *et al.*, US 5,807,718 .

The invention features novel nucleic acid molecules, including antisense oligonucleotides, siRNA and enzymatic nucleic acid molecules, and methods to modulate gene expression, for example, genes encoding K-Ras, H-Ras and/or N-Ras. In particular, the instant invention features nucleic-acid based molecules and methods to down-regulate the expression of K-Ras, H-Ras and/or N-Ras gene sequences.

The invention features one or more nucleic acid-based molecules and methods that independently or in combination modulate the expression of a gene or genes encoding Ras proteins. In particular embodiments, the invention features nucleic acid-based molecules and methods that modulate the expression of K-Ras gene, for example, Genbank Accession No. NM_004985; H-Ras gene, for example, Genbank Accession No. NM_005343; and/or N-Ras gene, for example, Genbank Accession No. NM_002524.

The description below of the various aspects and embodiments is provided with reference to exemplary K-Ras, H-Ras, and N-Ras genes, referred to hereinafter collectively as Ras. However, the various aspects and embodiments are directed to equivalent sequences and also to other genes which encode K-Ras, H-Ras and/or N-Ras proteins and similar proteins to K-Ras, H-Ras and/or N-Ras. For example, the invention relates to genes with homology to genes that encode K-Ras, H-Ras and/or N-Ras and genes that encode proteins with similar function to K-Ras, H-Ras, and N-Ras proteins. Those additional genes can be analyzed for target sites using the methods described herein. Thus, the modulation and the effects of such modulation of the other genes can be determined as described herein.

In one embodiment, the invention features the use of an enzymatic nucleic acid molecule, preferably in the hammerhead, NCH, G-cleaver, amberzyme, zinzyme and/or DNAzyme motif, to modulate the expression of a Ras gene or inhibit Ras activity. In one embodiment, the invention features the use of these enzymatic nucleic acid molecules to down-regulate the expression of a Ras gene or inhibit Ras activity. In another embodiment, the invention features the use of an antisense oligonucleotide molecule to modulate, for example, down-regulate, the expression of a Ras gene or inhibit Ras activity.

The invention features novel enzymatic nucleic acid molecules, siRNA molecules, and methods to modulate expression and/or activity of human immunodeficiency virus (HIV), for example HIV-1, HIV-2, and related viruses such as FIV-1 and SIV-1, or a HIV gene, for

example *LTR*, *nef*, *vif*, *tat*, or *rev*. In particular, the instant invention features nucleic-acid based molecules and methods to inhibit the replication of a HIV or related virus.

The invention features one or more nucleic acid-based molecules and methods that independently or in combination modulate the expression of gene(s) encoded by HIV and/or inhibit the replication of HIV. In particular embodiments, the invention features nucleic acid-based molecules and methods that modulate the expression of HIV-1 encoded genes, for example (Genbank Accession No. AJ302647); HIV-2 gene, for example (Genbank Accession No. NC_001722), FIV-1, for example (Genbank Accession No. NC_001482), SIV-1, for example (Genbank Accession No. M66437), *LTR*, for example included in (Genbank Accession No. AJ302647), *nef*, for example included in (Genbank Accession No. AJ302647), *vif*, for example included in (Genbank Accession No. AJ302647), *tat*, for example included in (Genbank Accession No. AJ302647), and *rev*, for example included in (Genbank Accession No. AJ302647).

The description below of the various aspects and embodiments is provided with reference to the exemplary HIV-1 gene, referred to herein as HIV. However, the various aspects and embodiments are also directed to other genes which encode HIV proteins and similar viruses to HIV. Those additional genes can be analyzed for target sites using the methods described for HIV. Thus, the inhibition and the effects of such inhibition of the other genes can be performed as described herein.

Due to the high sequence variability of the HIV genome, selection of nucleic acid molecules for broad therapeutic applications would likely involve the conserved regions of the HIV genome. Specifically, the present invention describes nucleic acid molecules that cleave the conserved regions of the HIV genome. Therefore, one nucleic acid molecule can be designed to cleave all the different isolates of HIV. Nucleic acid molecules designed against conserved regions of various HIV isolates can enable efficient inhibition of HIV replication in diverse subject populations and can ensure the effectiveness of the nucleic acid molecules against HIV quasi species which evolve due to mutations in the non-conserved regions of the HIV genome.

In one embodiment, the invention features the use of an enzymatic nucleic acid molecule, preferably in the hammerhead, NCH, G-cleaver, amberzyme, zinzyme and/or DNAzyme motif, to down-regulate the expression of HIV genes or inhibit the replication of HIV.

The invention features novel nucleic acid molecules, siRNA molecules and methods to modulate gene expression, for example, genes encoding HER2. In particular, the instant invention features nucleic-acid based molecules and methods to inhibit the expression of HER2.

5 The invention features one or more nucleic acid-based molecules and methods that independently or in combination modulate the expression of a gene or genes encoding HER2. In particular embodiments, the invention features nucleic acid-based molecules and methods that modulate the expression of HER2 gene, for example, Genbank Accession No. NM_004448.

10 The description below of the various aspects and embodiments is provided with reference to an exemplary HER2 gene, referred to herein as HER2 but also known as ERB2, ERB-B2, NEU, NGL, and v-ERB-B2. However, the various aspects and embodiments are also directed to other genes which encode HER2 proteins and similar proteins to HER2. Those additional genes can be analyzed for target sites using the methods described for
15 HER2. Thus, the inhibition and the effects of such inhibition of the other genes can be performed as described herein.

In one embodiment, the invention features the use of an enzymatic nucleic acid molecule, preferably in the hammerhead, NCH, G-cleaver, amberzyme, zinzyme and/or DNAzyme motif, to down-regulate the expression of HER2 genes or inhibit HER2 activity.

20 By "modulate" is meant that the expression of the gene, or level of RNAs or equivalent RNAs encoding one or more protein subunits or components, or activity of one or more proteins is up-regulated or down-regulated, such that the expression, level, or activity is greater than or less than that observed in the absence of the nucleic acid molecules of the invention.

25 By "inhibit" or "down-regulate" it is meant that the expression of the gene, or level of RNAs or equivalent RNAs encoding one or more protein subunits or components, or activity of one or more protein subunits or components, such as Ras, HIV, and/or HER2 protein or proteins, is reduced below that observed in the absence of the nucleic acid molecules of the invention. In one embodiment, inhibition or down-regulation with the enzymatic nucleic acid
30 molecule preferably is below that level observed in the presence of an enzymatically inactive or attenuated enzymatic nucleic acid molecule that is able to bind to the same site on the target RNA, but is unable to cleave that RNA. In another embodiment, inhibition or down-

regulation with an antisense oligonucleotide is preferably below that level observed in the presence of, for example, an oligonucleotide with scrambled sequence or with mismatches. In another embodiment, inhibition or down-regulation with an siRNA molecule is preferably below that level observed in the presence of, for example, an oligonucleotide with scrambled
5 sequence or with mismatches. In another embodiment, inhibition or down-regulation of Ras, HIV, or HER2 expression and/or activity with the nucleic acid molecule of the instant invention is greater in the presence of the nucleic acid molecule than in its absence.

By "up-regulate" is meant that the expression of the gene, or level of RNAs or equivalent RNAs encoding one or more protein subunits or components, or activity of one or
10 more protein subunits or components, such as Ras, HIV, or HER2 protein or proteins, is greater than that observed in the absence of the nucleic acid molecules of the invention. For example, the expression of a gene, such as Ras, HIV, or HER2 gene, can be increased in order to treat, prevent, ameliorate, or modulate a pathological condition caused or exacerbated by an absence or low level of gene expression.

By "enzymatic nucleic acid molecule" as used herein, is meant a nucleic acid molecule which has complementarity in a substrate binding region to a specified gene target, and also has an enzymatic activity which is active to specifically cleave target RNA. That is, the enzymatic nucleic acid molecule is able to intermolecularly cleave RNA and thereby
15 inactivate a target RNA molecule. These complementary regions allow sufficient hybridization of the enzymatic nucleic acid molecule to the target RNA and thus permit cleavage. One hundred percent complementarity is preferred, but complementarity as low as 50-75% can also be useful in this invention (see for example Werner and Uhlenbeck, 1995, *Nucleic Acids Research*, 23, 2092-2096; Hammann *et al.*, 1999, *Antisense and Nucleic Acid Drug Dev.*, 9, 25-31). The nucleic acids can be modified at the base, sugar, and/or phosphate
20 groups. The term DNAzyme-based enzymatic nucleic acid is used interchangeably with phrases such as catalytic DNA, aptazyme or aptamer-binding DNAzyme, regulatable DNAzyme, catalytic oligonucleotides, nucleozyme, DNAzyme, endoribonuclease, endonuclease, minizyme, leadzyme, oligozyme or DNA enzyme. All of these terminologies describe nucleic acid molecules with enzymatic activity. The specific enzymatic nucleic acid
25 molecules described in the instant application are not limiting in the invention and those skilled in the art will recognize that all that is important in an enzymatic nucleic acid molecule of this invention is that it have a specific substrate binding site which is complementary to one or more of the target nucleic acid regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart a nucleic acid
30 cleaving and/or ligation activity to the molecule.
35

By "nucleic acid molecule" as used herein is meant a molecule having nucleotides. The nucleic acid can be single, double, or multiple stranded and can comprise modified or unmodified nucleotides or non-nucleotides or various mixtures and combinations thereof.

5 By "enzymatic portion" or "catalytic domain" is meant that portion/region of the enzymatic nucleic acid molecule essential for cleavage of a nucleic acid substrate (for example see Figures 1-4).

10 By "substrate binding arm" or "substrate binding domain" is meant that portion/region of a enzymatic nucleic acid which is able to interact, for example via complementarity (*i.e.*, able to base-pair with), with a portion of its substrate. Preferably, such complementarity is 100%, but can be less if desired. For example, as few as 10 bases out of 14 can be base-paired (see for example Werner and Uhlenbeck, 1995, *Nucleic Acids Research*, 23, 2092-2096; Hammann *et al.*, 1999, *Antisense and Nucleic Acid Drug Dev.*, 9, 25-31). Examples of such arms are shown generally in Figures 1-3. That is, these arms contain sequences within a enzymatic nucleic acid which are intended to bring enzymatic nucleic acid and target RNA
15 together through complementary base-pairing interactions. The enzymatic nucleic acid of the invention can have binding arms that are contiguous or non-contiguous and can be of varying lengths. The length of the binding arm(s) are preferably greater than or equal to four nucleotides and of sufficient length to stably interact with the target RNA; preferably 12-100 nucleotides; more preferably 14-24 nucleotides long (see for example Werner and Uhlenbeck, 20 *supra*; Hamman *et al.*, *supra*; Hampel *et al.*, EP0360257; Berzal-Herranz *et al.*, 1993, *EMBO J.*, 12, 2567-73). If two binding arms are chosen, the design is such that the length of the binding arms are symmetrical (*i.e.*, each of the binding arms is of the same length; *e.g.*, five and five nucleotides, or six and six nucleotides, or seven and seven nucleotides long) or asymmetrical (*i.e.*, the binding arms are of different length; *e.g.*, six and three nucleotides; 25 three and six nucleotides long; four and five nucleotides long; four and six nucleotides long; four and seven nucleotides long; and the like).

By "Inozyme" or "NCH" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described as NCH Rz in Figure 1 and in Ludwig *et al.*, International PCT Publication No. WO 98/58058 and US Patent Application Serial No. 30 08/878,640. Inozymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NCH/, where N is a nucleotide, C is cytidine and H is adenosine, uridine or cytidine, and "/" represents the cleavage site. H is used interchangeably with X. Inozymes can also possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NCN/, where N is a nucleotide, C is cytidine, and "/" represents the cleavage site. "T"

in Figure 1 represents an Inosine nucleotide, preferably a ribo-Inosine or xylo-Inosine nucleoside.

By "G-cleaver" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described as G-cleaver Rz in Figure 1 and in Eckstein *et al.*, US 6,127,173. G-cleavers possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NYN/, where N is a nucleotide, Y is uridine or cytidine and "/" represents the cleavage site. G-cleavers can be chemically modified as is generally shown in Figure 1.

By "amberzyme" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described in Figure 2 and in Beigelman *et al.*, International PCT publication No. WO 99/55857 and US Patent Application Serial No. 09/476,387. Amberzymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NG/N, where N is a nucleotide, G is guanosine, and "/" represents the cleavage site. Amberzymes can be chemically modified to increase nuclease stability through substitutions as are generally shown in Figure 2. In addition, differing nucleoside and/or non-nucleoside linkers can be used to substitute the 5'-gaaa-3' loops shown in the figure. Amberzymes represent a non-limiting example of an enzymatic nucleic acid molecule that does not require a ribonucleotide (2'-OH) group within its own nucleic acid sequence for activity.

By "zinzyme" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described in Figure 3 and in Beigelman *et al.*, International PCT publication No. WO 99/55857 and US Patent Application Serial No. 09/918,728. Zinzymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet including but not limited to YG/Y, where Y is uridine or cytidine, and G is guanosine and "/" represents the cleavage site. Zinzymes can be chemically modified to increase nuclease stability through substitutions as are generally shown in Figure 3, including substituting 2'-O-methyl guanosine nucleotides for guanosine nucleotides. In addition, differing nucleotide and/or non-nucleotide linkers can be used to substitute the 5'-gaaa-2' loop shown in the figure. Zinzymes represent a non-limiting example of an enzymatic nucleic acid molecule that does not require a ribonucleotide (2'-OH) group within its own nucleic acid sequence for activity.

By 'DNAzyme' is meant, an enzymatic nucleic acid molecule that does not require the presence of a 2'-OH group within its own nucleic acid sequence for activity. In particular

embodiments the enzymatic nucleic acid molecule can have an attached linker or linkers or other attached or associated groups, moieties, or chains containing one or more nucleotides with 2'-OH groups. DNAzymes can be synthesized chemically or expressed endogenously *in vivo*, by means of a single stranded DNA vector or equivalent thereof. An example of a DNAzyme is shown in **Figure 4** and is generally reviewed in Usman *et al.*, US patent No., 6,159,714; Chartrand *et al.*, 1995, *NAR* 23, 4092; Breaker *et al.*, 1995, *Chem. Bio.* 2, 655; Santoro *et al.*, 1997, *PNAS* 94, 4262; Breaker, 1999, *Nature Biotechnology*, 17, 422-423; and Santoro *et al.*, 2000, *J. Am. Chem. Soc.*, 122, 2433-39. The "10-23" DNAzyme motif is one particular type of DNAzyme that was evolved using *in vitro* selection, see Santoro *et al.*, *supra* and as generally described in Joyce *et al.*, US 5,807,718. Additional DNAzyme motifs can be selected by using techniques similar to those described in these references, and hence, are within the scope of the present invention. DNAzymes of the invention can comprise nucleotides modified at the nucleic acid base, sugar, or phosphate backbone. Non-limiting examples of sugar modifications that can be used in DNAzymes of the invention include 2'-O-alkyl modifications such as 2'-O-methyl or 2'-O-allyl, 2'-C-alkyl modifications such as 2'-C-allyl, 2'-deoxy-2'-amino, 2'-halo modifications such as 2'-fluoro, 2'-chloro, or 2'-bromo, isomeric modifications such as arabinofuranose or xylofuranose based nucleic acids, and other sugar modifications such as 4'-thio or 4'-carbocyclic nucleic acids. Non-limiting examples of nucleic acid based modifications that can be used in DNAzymes of the invention include modified purine heterocycles, G-clamp heterocycles, and various modified pyrimidine cycles. Non-limiting examples of backbone modifications that can be used in DNAzymes of the invention include phosphorothioate, phosphorodithioate, phosphoramidate, and methylphosphonate internucleotide linkages. DNAzymes of the invention can comprise naturally occurring nucleic acids, chimeras of chemically modified and naturally occurring nucleic acids, or completely modified nucleic acids.

In general, enzymatic nucleic acids act by first binding to a target RNA. Such binding occurs through the target binding portion of a enzymatic nucleic acid that is held in close proximity to an enzymatic portion of the molecule that acts to cleave the target RNA. Thus, the enzymatic nucleic acid first recognizes and then binds a target RNA through complementary base-pairing, and once bound to the correct site, acts enzymatically to cut the target RNA. Strategic cleavage of such a target RNA will destroy its ability to direct synthesis of an encoded protein. After an enzymatic nucleic acid has bound and cleaved its RNA target, it is released from that RNA to search for another target and can repeatedly bind and cleave new targets. Thus, a single enzymatic nucleic acid molecule is able to cleave many molecules of target RNA. In addition, the enzymatic nucleic acid molecule is a highly

specific inhibitor of gene expression, with the specificity of inhibition depending not only on the base-pairing mechanism of binding to the target RNA, but also on the mechanism of target RNA cleavage. Single mismatches, or base-substitutions, near the site of cleavage can completely eliminate catalytic activity of an enzymatic nucleic acid molecule.

5 By "sufficient length" is meant an oligonucleotide of greater than or equal to 3 nucleotides that is of a length great enough to provide the intended function under the expected condition. For example, for binding arms of enzymatic nucleic acid "sufficient length" means that the binding arm sequence is long enough to provide stable binding to a target site under the expected binding conditions. Preferably, the binding arms are not so
10 long as to prevent useful turnover of the nucleic acid molecule.

By "stably interact" is meant interaction of oligonucleotides with target nucleic acid molecules (*e.g.*, by forming hydrogen bonds with complementary nucleotides in the target under physiological conditions) that is sufficient to the intended purpose (*e.g.*, cleavage of target RNA by an enzyme).

15 By "equivalent" RNA to Ras is meant to include those naturally occurring RNA molecules having homology (partial or complete) to Ras nucleic acids or encoding for proteins with similar function as Ras proteins in various organisms, including humans, rodents, primates, rabbits, pigs, protozoans, fungi, plants, and other microorganisms and parasites. The equivalent RNA sequence can also include, in addition to the coding region,
20 regions such as a 5'-untranslated region, a 3'-untranslated region, introns, a intron-exon junction and the like.

By "equivalent" RNA to HIV is meant to include those naturally occurring RNA molecules having homology (partial or complete) to HIV nucleic acids or encoding for proteins with similar function as HIV proteins in various organisms, including human, rodent,
25 primate, rabbit, pig, protozoans, fungi, plants, and other microorganisms and parasites. The equivalent RNA sequence also includes in addition to the coding region, regions such as 5'-untranslated region, 3'-untranslated region, introns, intron-exon junction and the like.

By "equivalent" RNA to HER2 is meant to include those naturally occurring RNA molecules having homology (partial or complete) to HER2 nucleic acids or encoding for
30 proteins with similar function as HER2 proteins in various organisms, including humans, rodents, primates, rabbits, pigs, protozoans, fungi, plants, and other microorganisms and parasites. The equivalent RNA sequence also includes, in addition to the coding region,

regions such as a 5'-untranslated region, a 3'-untranslated region, introns, a intron-exon junction and the like.

By "homology" is meant the nucleotide sequence of two or more nucleic acid molecules is partially or completely identical.

5 By "component" of HIV is meant a peptide or protein expressed from an HIV gene, for example *nef*, *vif*, *tat*, or *rev* viral gene products.

By "component" of HER2 is meant a peptide or protein subunit expressed from a HER2 gene.

10 By "component" of Ras is meant a peptide or protein subunit expressed from a Ras gene.

By "gene" it is meant a nucleic acid that encodes an RNA, for example, nucleic acid sequences including but not limited to structural genes encoding a polypeptide.

"Complementarity" refers to the ability of a nucleic acid to form hydrogen bond or bonds with another RNA sequence by either traditional Watson-Crick or other non-traditional
15 types. In reference to the nucleic molecules of the present invention, the binding free energy for a nucleic acid molecule with its target or complementary sequence is sufficient to allow the relevant function of the nucleic acid to proceed, *e.g.*, enzymatic nucleic acid cleavage, antisense or triple helix inhibition. Determination of binding free energies for nucleic acid molecules is well known in the art (see, *e.g.*, Turner *et al.*, 1987, *CSH Symp. Quant. Biol.* LII
20 pp.123-133; Frier *et al.*, 1986, *Proc. Nat. Acad. Sci. USA* 83:9373-9377; Turner *et al.*, 1987, *J. Am. Chem. Soc.* 109:3783-3785). A percent complementarity indicates the percentage of contiguous residues in a nucleic acid molecule that can form hydrogen bonds (*e.g.*, Watson-Crick base pairing) with a second nucleic acid sequence (*e.g.*, 5, 6, 7, 8, 9, 10 out of 10 being 50%, 60%, 70%, 80%, 90%, and 100% complementary). "Perfectly complementary" means
25 that all the contiguous residues of a nucleic acid sequence will hydrogen bond with the same number of contiguous residues in a second nucleic acid sequence.

By "RNA" is meant a molecule comprising at least one ribonucleotide residue. By "ribonucleotide" or "2'-OH" is meant a nucleotide with a hydroxyl group at the 2' position of a β -D-ribo-furanose moiety.

30 By "decoy " is meant a nucleic acid molecule, for example RNA or DNA, or aptamer that is designed to preferentially bind to a predetermined ligand. Such binding can result in

the inhibition or activation of a target molecule. A decoy or aptamer can compete with a naturally occurring binding target for the binding of a specific ligand. For example, it has been shown that over-expression of HIV trans-activation response (TAR) RNA can act as a "decoy" and efficiently binds HIV tat protein, thereby preventing it from binding to TAR sequences encoded in the HIV RNA (Sullenger *et al.*, 1990, *Cell*, 63, 601-608). This is but a specific example and those in the art will recognize that other embodiments can be readily generated using techniques generally known in the art, see for example Gold *et al.*, 1995, *Annu. Rev. Biochem.*, 64, 763; Brody and Gold, 2000, *J. Biotechnol.*, 74, 5; Sun, 2000, *Curr. Opin. Mol. Ther.*, 2, 100; Kusser, 2000, *J. Biotechnol.*, 74, 27; Hermann and Patel, 2000, *Science*, 287, 820; and Jayasena, 1999, *Clinical Chemistry*, 45, 1628. Similarly, a decoy can be designed to bind to Ras and block the binding of Ras or a decoy can be designed to bind to Ras and prevent interaction with the Ras protein.

By "aptamer" or "nucleic acid aptamer" as used herein is meant a nucleic acid molecule that binds specifically to a target molecule wherein the nucleic acid molecule has sequence that is distinct from sequence recognized by the target molecule in its natural setting. Alternately, an aptamer can be a nucleic acid molecule that binds to a target molecule where the target molecule does not naturally bind to a nucleic acid. The target molecule can be any molecule of interest. For example, the aptamer can be used to bind to a ligand binding domain of a protein, thereby preventing interaction of the naturally occurring ligand with the protein. Similarly, the nucleic acid molecules of the instant invention can bind to RAS, Her-2 or HIV encoded RNA or proteins receptors to block activity of the activity of target protein or nucleic acid. This is a non-limiting example and those in the art will recognize that other embodiments can be readily generated using techniques generally known in the art, see for example Gold *et al.*, US 5,475,096 and 5,270,163; Gold *et al.*, 1995, *Annu. Rev. Biochem.*, 64, 763; Brody and Gold, 2000, *J. Biotechnol.*, 74, 5; Sun, 2000, *Curr. Opin. Mol. Ther.*, 2, 100; Kusser, 2000, *J. Biotechnol.*, 74, 27; Hermann and Patel, 2000, *Science*, 287, 820; and Jayasena, 1999, *Clinical Chemistry*, 45, 1628.

The term "short interfering RNA" or "siRNA" as used herein refers to a double stranded nucleic acid molecule capable of RNA interference "RNAi", see for example Bass, 2001, *Nature*, 411, 428-429; Elbashir *et al.*, 2001, *Nature*, 411, 494-498; and Kreutzer *et al.*, International PCT Publication No. WO 00/44895; Zernicka-Goetz *et al.*, International PCT Publication No. WO 01/36646; Fire, International PCT Publication No. WO 99/32619; Plaetinck *et al.*, International PCT Publication No. WO 00/01846; Mello and Fire, International PCT Publication No. WO 01/29058; Deschamps-Depaillette, International PCT Publication No. WO 99/07409; and Li *et al.*, International PCT Publication No. WO

00/44914. As used herein, siRNA molecules need not be limited to those molecules containing only RNA, but further encompasses chemically modified nucleotides and non-nucleotides.

5 Nucleic acid molecules that modulate expression of Ras-specific RNAs represent a therapeutic approach to treat cancer, including, but not limited to colorectal cancer, bladder cancer, lung cancer, pancreatic cancer, breast cancer, or prostate cancer and any other cancer, disease or condition that responds to the modulation of Ras expression.

10 Nucleic acid molecules that modulate expression of HIV-specific RNAs also represent a therapeutic approach to treat acquired immunodeficiency syndrome (AIDS) and/or any other disease, condition, or syndrome which respond to the modulation of HIV expression.

Nucleic acid molecules that modulate expression of HER2-specific RNAs represent a therapeutic approach to treat cancer, including, but not limited to breast and ovarian cancer and any other cancer, disease or condition that responds to the modulation of HER2 expression.

15 In one embodiment of the inventions described herein, the enzymatic nucleic acid molecule is formed in a hammerhead or hairpin motif, but can also be formed in the motif of a hepatitis delta virus, group I intron, group II intron or RNase P RNA (in association with an RNA guide sequence), *Neurospora* VS RNA, DNAzymes, NCH cleaving motifs, or G-cleavers. Examples of such hammerhead motifs are described by Dreyfus, *supra*, Rossi *et al.*,
20 1992, *AIDS Research and Human Retroviruses* 8, 183; of hairpin motifs by Hampel *et al.*, EP0360257, Hampel and Tritz, 1989 *Biochemistry* 28, 4929, Feldstein *et al.*, 1989, *Gene* 82, 53, Haseloff and Gerlach, 1989, *Gene*, 82, 43, and Hampel *et al.*, 1990 *Nucleic Acids Res.* 18, 299; Chowrira & McSwiggen, US. Patent No. 5,631,359; of the hepatitis delta virus motif is described by Perrotta and Been, 1992 *Biochemistry* 31, 16; of the RNase P motif by Guerrier-Takada *et al.*, 1983 *Cell* 35, 849; Forster and Altman, 1990, *Science* 249, 783; Li and Altman, 25 1996, *Nucleic Acids Res.* 24, 835; *Neurospora* VS RNA ribozyme motif is described by Collins (Saville and Collins, 1990 *Cell* 61, 685-696; Saville and Collins, 1991 *Proc. Natl. Acad. Sci. USA* 88, 8826-8830; Collins and Olive, 1993 *Biochemistry* 32, 2795-2799; Guo and Collins, 1995, *EMBO. J.* 14, 363); Group II introns are described by Griffin *et al.*, 1995,
30 *Chem. Biol.* 2, 761; Michels and Pyle, 1995, *Biochemistry* 34, 2965; Pyle *et al.*, International PCT Publication No. WO 96/22689; of the Group I intron by Cech *et al.*, U.S. Patent 4,987,071 and of DNAzymes by Usman *et al.*, International PCT Publication No. WO 95/11304; Chartrand *et al.*, 1995, *NAR* 23, 4092; Breaker *et al.*, 1995, *Chem. Bio.* 2, 655; Santoro *et al.*, 1997, *PNAS* 94, 4262, and Beigelman *et al.*, International PCT publication No.

WO 99/55857. NCH cleaving motifs are described in Ludwig & Sproat, International PCT Publication No. WO 98/58058; and G-cleavers are described in Kore *et al.*, 1998, *Nucleic Acids Research* 26, 4116-4120 and Eckstein *et al.*, International PCT Publication No. WO 99/16871. Additional motifs such as the Aptazyme (Breaker *et al.*, WO 98/43993),
5 Amberzyme (Class I motif; Figure 2; Beigelman *et al.*, U.S. Serial No. 09/301,511) and Zinzyme (Figure 3) (Beigelman *et al.*, U.S. Serial No. 09/301,511), all included by reference herein including drawings, can also be used in the present invention. These specific motifs or configurations are not limiting in the invention and those skilled in the art will recognize that all that is important in an enzymatic nucleic acid molecule of this invention is that it has a
10 specific substrate binding site which is complementary to one or more of the target gene RNA regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart an RNA cleaving activity to the molecule (Cech *et al.*, U.S. Patent No. 4,987,071).

In one embodiment of the present invention, a nucleic acid molecule of the instant
15 invention can be between about 10 and 100 nucleotides in length. Exemplary enzymatic nucleic acid molecules of the invention are shown in the Tables herein. For example, enzymatic nucleic acid molecules of the invention are preferably between about 15 and 50 nucleotides in length, more preferably between about 25 and 40 nucleotides in length, *e.g.*, 34, 36, or 38 nucleotides in length (for example see Jarvis *et al.*, 1996, *J. Biol. Chem.*, 271, 29107-29112). Exemplary DNAzymes of the invention are preferably between about 15 and
20 40 nucleotides in length, more preferably between about 25 and 35 nucleotides in length, *e.g.*, 29, 30, 31, or 32 nucleotides in length (see for example Santoro *et al.*, 1998, *Biochemistry*, 37, 13330-13342; Chartrand *et al.*, 1995, *Nucleic Acids Research*, 23, 4092-4096). Exemplary antisense molecules of the invention are preferably between about 15 and 75
25 nucleotides in length, more preferably between about 20 and 35 nucleotides in length, *e.g.*, 25, 26, 27, or 28 nucleotides in length (see for example Woolf *et al.*, 1992, *PNAS*, 89, 7305-7309; Milner *et al.*, 1997, *Nature Biotechnology*, 15, 537-541). Exemplary triplex forming oligonucleotide molecules of the invention are preferably between about 10 and 40 nucleotides in length, more preferably between about 12 and 25 nucleotides in length, *e.g.*,
30 18, 19, 20, or 21 nucleotides in length (see for example Maher *et al.*, 1990, *Biochemistry*, 29, 8820-8826; Strobel and Dervan, 1990, *Science*, 249, 73-75). Those skilled in the art will recognize that all that is required is for a nucleic acid molecule to be of length and conformation sufficient and suitable for the nucleic acid molecule to interact with its target and/or catalyze a reaction contemplated herein. The length of nucleic acid molecules of the
35 instant invention are not limiting within the general limits stated.

Preferably, a nucleic acid molecule that modulates, for example, down-regulates Ras, HIV, and/or HER2 expression and/or activity, comprises between 12 and 100 bases complementary to a RNA molecule of Ras, HIV, and/or HER2 respectively. Even more preferably, a nucleic acid molecule that modulates Ras, HIV, and/or HER2 expression
5 comprises between 14 and 24 bases complementary to a RNA molecule of Ras, HIV, and/or HER2 respectively.

The invention provides a method for producing a class of nucleic acid-based gene modulating agents that exhibit a high degree of specificity for RNA of a desired target. For example, an enzymatic nucleic acid molecule is preferably targeted to a highly conserved
10 sequence region of target RNAs encoding Ras (and specifically a Ras gene) such that specific treatment of a disease or condition can be provided with either one or several nucleic acid molecules of the invention. Such nucleic acid molecules can be delivered exogenously to specific tissue or cellular targets as required. Alternatively, the nucleic acid molecules (e.g., enzymatic nucleic acid molecules, siRNA, antisense, and/or DNazymes) can be expressed
15 from DNA and/or RNA vectors that are delivered to specific cells.

As used herein "cell" is used in its usual biological sense, and does not refer to an entire multicellular organism. A cell can, for example, be *in vitro*, e.g., in cell culture, or present in a multicellular organism, including, e.g., birds, plants and mammals such as humans, cows, sheep, apes, monkeys, swine, dogs, and cats. The cell can be prokaryotic (e.g., bacterial cell)
20 or eukaryotic (e.g., mammalian or plant cell).

By "Ras proteins" is meant, a peptide or protein comprising Ras tyrosine kinase-type cell surface receptor or a peptide or protein encoded by a Ras gene, such as K-Ras, H-Ras, or N-Ras.

By "HIV proteins" is meant, a peptide or protein comprising a component of HIV or a
25 peptide or protein encoded by a HIV gene.

By "HER2 proteins" is meant, a peptide or protein comprising HER2/ERB2/NEU tyrosine kinase-type cell surface receptor or a peptide or protein encoded by a HER2/ERB2/NEU gene.

By "highly conserved sequence region" is meant, a nucleotide sequence of one or more
30 regions in a target gene that does not vary significantly from one generation to the other or from one biological system to the other.

Nucleic acid-based modulators, including inhibitors, of Ras expression are useful for the prevention and/or treatment of cancer, including but not limited to breast cancer and ovarian cancer and any other disease or condition that respond to the modulation of Ras expression.

5 Nucleic acid-based inhibitors of HIV expression are useful for the prevention and/or treatment of acquired immunodeficiency disease (AIDS) and related diseases and conditions, including but not limited to Kaposi's sarcoma, lymphoma, cervical cancer, squamous cell carcinoma, cardiac myopathy, rheumatic diseases, and opportunistic infection, for example
10 Pneumocystis carinii, Cytomegalovirus, Herpes simplex, Mycobacteria, Cryptococcus, Toxoplasma, Progressive multifocal leucoencephalopathy (Papovavirus), Mycobacteria, Aspergillus, Cryptococcus, Candida, Cryptosporidium, Isospora belli, Microsporidia and any other disease or condition which respond to the modulation of HIV expression.

Nucleic acid-based inhibitors of HER2 expression are useful for the prevention and/or treatment of cancer, including but not limited to breast cancer and ovarian cancer and any
15 other disease or condition that respond to the modulation of HER2 expression.

By "related" is meant that the reduction of RAS, HIV, or HER2 expression (specifically RAS, HIV, or HER2 genes respectively) RNA levels and thus reduction in the level of the respective protein relieves, to some extent, the symptoms of the disease or condition.

The nucleic acid-based molecules of the invention can be added directly, or can be
20 complexed with cationic lipids, packaged within liposomes, or otherwise delivered to target cells or tissues. The nucleic acid or nucleic acid complexes can be locally administered to relevant tissues *ex vivo*, or *in vivo* through injection or infusion pump, with or without their incorporation in biopolymers. In certain embodiments, the enzymatic nucleic acid molecules comprise sequences that are complementary to the substrate sequences in the Tables herein.
25 Examples of such enzymatic nucleic acid molecules also are shown in the Tables herein. Examples of such enzymatic nucleic acid molecules consist essentially of sequences defined in these tables.

In another embodiment, the invention features siRNA, antisense nucleic acid molecules and 2-5A chimeras comprising sequences complementary to the substrate sequences shown in
30 the Tables herein. Such nucleic acid molecules can comprise sequences as shown for the binding arms of the enzymatic nucleic acid molecules in the Tables. Similarly, triplex molecules can be targeted to corresponding DNA target regions; such molecules can comprise the DNA equivalent of a target sequence or a sequence complementary to the specified target

(substrate) sequence. Typically, antisense molecules are complementary to a target sequence along a single contiguous sequence of the antisense molecule. However, in certain embodiments, an antisense molecule can bind to a substrate such that the substrate molecule forms a loop, and/or an antisense molecule can bind such that the antisense molecule forms a loop. Thus, the antisense molecule can be complementary to two or more non-contiguous substrate sequences. In addition, two or more non-contiguous sequence portions of an antisense molecule can be complementary to a target sequence.

By "consists essentially of" is meant that the active nucleic acid molecule of the invention, for example, an enzymatic nucleic acid molecule, contains an enzymatic center or core equivalent to those in the examples, and binding arms able to bind RNA such that cleavage at the target site occurs. Other sequences can be present that do not interfere with such cleavage. Thus, a core region of an enzymatic nucleic acid molecule can, for example, include one or more loop, stem-loop structure, or linker that does not prevent enzymatic activity. Thus, various regions in the sequences in the Tables can be such a loop, stem-loop, nucleotide linker, and/or non-nucleotide linker and can be represented generally as sequence "X". The nucleic acid molecules of the instant invention, such as Hammerhead, Inozyme, G-cleaver, amberzyme, zinzyme, DNAzyme, antisense, 2-5A antisense, triplex forming nucleic acid, and decoy nucleic acids, can contain other sequences or non-nucleotide linkers that do not interfere with the function of the nucleic acid molecule.

Sequence X can be a linker of ≥ 2 nucleotides in length, preferably 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 26, 30, where the nucleotides can preferably be internally base-paired to form a stem of preferably ≥ 2 base pairs. Alternatively or in addition, sequence X can be a non-nucleotide linker. In yet another embodiment, the nucleotide linker X can be a nucleic acid aptamer, such as an ATP aptamer, Ras Rev aptamer (RRE), Ras Tat aptamer (TAR) and others (for a review see Gold *et al.*, 1995, *Annu. Rev. Biochem.*, 64, 763; and Szostak & Ellington, 1993, in *The RNA World*, ed. Gesteland and Atkins, pp. 511, CSH Laboratory Press). A "nucleic acid aptamer" as used herein is meant to indicate a nucleic acid sequence capable of interacting with a ligand. The ligand can be any natural or a synthetic molecule, including but not limited to a resin, metabolites, nucleosides, nucleotides, drugs, toxins, transition state analogs, peptides, lipids, proteins, amino acids, nucleic acid molecules, hormones, carbohydrates, receptors, cells, viruses, bacteria and others.

In yet another embodiment, a non-nucleotide linker X is as defined herein. Non-nucleotides as can include abasic nucleotide, polyether, polyamine, polyamide, peptide, carbohydrate, lipid, or polyhydrocarbon compounds. Specific examples include those

described by Seela and Kaiser, *Nucleic Acids Res.* 1990, 18:6353 and *Nucleic Acids Res.* 1987, 15:3113; Cload and Schepartz, *J. Am. Chem. Soc.* 1991, 113:6324; Richardson and Schepartz, *J. Am. Chem. Soc.* 1991, 113:5109; Ma *et al.*, *Nucleic Acids Res.* 1993, 21:2585 and *Biochemistry* 1993, 32:1751; Durand *et al.*, *Nucleic Acids Res.* 1990, 18:6353; McCurdy
5 *et al.*, *Nucleosides & Nucleotides* 1991, 10:287; Jsche *et al.*, *Tetrahedron Lett.* 1993, 34:301; Ono *et al.*, *Biochemistry* 1991, 30:9914; Arnold *et al.*, International Publication No. WO 89/02439; Usman *et al.*, International Publication No. WO 95/06731; Dudycz *et al.*, International Publication No. WO 95/11910 and Ferentz and Verdine, *J. Am. Chem. Soc.* 1991, 113:4000, all hereby incorporated by reference herein. A "non-nucleotide" further
10 means any group or compound that can be incorporated into a nucleic acid chain in the place of one or more nucleotide units, including either sugar and/or phosphate substitutions, and allows the remaining bases to exhibit their enzymatic activity. The group or compound can be abasic in that it does not contain a commonly recognized nucleotide base, such as adenosine, guanine, cytosine, uracil or thymine. Thus, in a preferred embodiment, the
15 invention features an enzymatic nucleic acid molecule having one or more non-nucleotide moieties, and having enzymatic activity to cleave an RNA or DNA molecule.

In another aspect of the invention, enzymatic nucleic acid molecules, siRNA molecules or antisense molecules that interact with target RNA molecules and modulate gene expression activity are expressed from transcription units inserted into DNA or RNA vectors. The
20 recombinant vectors are preferably DNA plasmids or viral vectors. Enzymatic nucleic acid molecule or antisense expressing viral vectors can be constructed based on, but not limited to, adeno-associated virus, retrovirus, adenovirus, or alphavirus as well as others known in the art. Preferably, recombinant vectors capable of expressing enzymatic nucleic acid molecules or antisense are delivered as described below, and persist in target cells. Alternatively, viral
25 vectors can be used that provide for transient expression of enzymatic nucleic acid molecules or antisense. Such vectors can be repeatedly administered as necessary. Once expressed, the enzymatic nucleic acid molecules or antisense bind to target RNA and modulate its function or expression. Delivery of enzymatic nucleic acid molecule or antisense expressing vectors can be systemic, such as by intravenous or intramuscular administration, by administration to
30 target cells ex-planted from the patient followed by reintroduction into the patient, or by any other means that allows for introduction into a desired target cell. Antisense DNA and DNazymes can be expressed via the use of a single stranded DNA intracellular expression vector.

By "vectors" is meant any nucleic acid- and/or viral-based technique used to deliver a
35 desired nucleic acid.

By "subject" or "patient" is meant an organism that is a donor or recipient of explanted cells or the cells of the organism. "Subject" or "patient" also refers to an organism to which the nucleic acid molecules of the invention can be administered. Preferably, a subject or patient is a mammal or mammalian cells. More preferably, a subject or patient is a human or human cells.

By "enhanced enzymatic activity" is meant to include activity measured in cells and/or *in vivo* where the activity is a reflection of both the catalytic activity and the stability of the nucleic acid molecules of the invention. In this invention, the product of these properties can be increased *in vivo* compared to an all RNA enzymatic nucleic acid or all DNA enzyme, for example, with a nucleic acid molecule comprising chemical modifications. In some cases, the activity or stability of the nucleic acid molecule can be decreased (i.e., less than ten-fold), but the overall activity of the nucleic acid molecule is enhanced, *in vivo*.

Nucleic acid molecules of the instant invention, individually, or in combination or in conjunction with other drugs, can be used to treat diseases or conditions discussed above. For example, to treat a disease or condition associated with the levels of Ras, HIV, or HER2, a subject can be treated, or other appropriate cells can be treated, as is evident to those skilled in the art, individually or in combination with one or more drugs under conditions suitable for the treatment.

In a further embodiment, the described molecules, such as antisense, siRNA, or enzymatic nucleic acid molecules, can be used in combination with other known treatments to treat conditions or diseases discussed above. For example, the described molecules can be used in combination with one or more known therapeutic agents to treat cancer, for example colorectal cancer, bladder cancer, lung cancer, pancreatic cancer, breast cancer, or prostate cancer, and any other disease or condition that respond to the modulation of Ras expression.

In another embodiment, the invention features nucleic acid-based inhibitors (e.g., enzymatic nucleic acid molecules, (including DNazymes), siRNA and methods for their use to down regulate or inhibit the expression of genes (e.g., Ras genes) capable of progression and/or maintenance of cancer and/or other disease states that respond to the modulation of Ras expression.

In a further embodiment, the described molecules, such as antisense, siRNA, or enzymatic nucleic acids, can be used in combination with other known treatments to treat conditions or diseases discussed above. For example, the described molecules can be used in combination with one or more known therapeutic agents to treat acquired immunodeficiency

disease (AIDS) and related diseases and conditions, including but not limited to Kaposi's sarcoma, lymphoma, cervical cancer, squamous cell carcinoma, cardiac myopathy, rheumatic diseases, and opportunistic infection, for example Pneumocystis carinii, Cytomegalovirus, Herpes simplex, Mycobacteria, Cryptococcus, Toxoplasma, Progressive multifocal
5 leucoencephalopathy (Papovavirus), Mycobacteria, Aspergillus, Cryptococcus, Candida, Cryptosporidium, Isospora belli, Microsporidia and any other disease or condition which respond to the modulation of HIV expression.

Nucleic acid molecules of the instant invention, individually, or in combination or in conjunction with other drugs, can be used to treat diseases or conditions discussed above. For
10 example, to treat a disease or condition associated with the levels of HER2, a patient can be treated, or other appropriate cells can be treated, as is evident to those skilled in the art, individually or in combination with one or more drugs under conditions suitable for the treatment.

In a further embodiment, the described molecules, such as antisense, siRNA or enzymatic nucleic acid molecules, can be used in combination with other known treatments to
15 treat conditions or diseases discussed above. For example, the described molecules can be used in combination with one or more known therapeutic agents to treat cancer, for example ovarian cancer and/or breast cancer, and any other disease or condition that respond to the modulation of HER2 expression.

In another embodiment, the invention features nucleic acid-based inhibitors (e.g., enzymatic nucleic acid molecules, (including ribozymes, antisense nucleic acids, 2-5A antisense chimeras, triplex DNA, antisense nucleic acids containing RNA cleaving chemical
20 groups), siRNA and methods for their use to down regulate or inhibit the expression of genes (e.g., HER2 genes) capable of progression and/or maintenance of cancer and/or other disease states that respond to the modulation of HER2 expression.
25

By "comprising" is meant including, but not limited to, whatever follows the word "comprising". Thus, use of the term "comprising" indicates that the listed elements are required or mandatory, but that other elements are optional and may or may not be present. By "consisting of" is meant including, and limited to, whatever follows the phrase "consisting
30 of".

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

Mechanism of action of Nucleic Acid Molecules of the Invention as is Known in the Art

Antisense: Antisense molecules can be modified or unmodified RNA, DNA, or mixed polymer oligonucleotides and primarily function by specifically binding to matching sequences resulting in inhibition of peptide synthesis (Wu-Pong, Nov 1994, *BioPharm*, 20-33). The antisense oligonucleotide binds to target RNA by Watson Crick base-pairing and blocks gene expression by preventing ribosomal translation of the bound sequences either by steric blocking or by activating RNase H enzyme. Antisense molecules can also alter protein synthesis by interfering with RNA processing or transport from the nucleus into the cytoplasm (Mukhopadhyay & Roth, 1996, *Crit. Rev. in Oncogenesis* 7, 151-190).

In addition, binding of single stranded DNA to RNA can result in nuclease degradation of the heteroduplex (Wu-Pong, *supra*; Crooke, *supra*). Backbone modified DNA chemistry which have been thus far been shown to act as substrates for RNase H are phosphorothioates, phosphorodithioates, and borontrifluoridates. In addition, 2'-arabino and 2'-fluoro arabino-containing oligos can also activate RNase H activity.

A number of antisense molecules have been described that utilize novel configurations of chemically modified nucleotides, secondary structure, and/or RNase H substrate domains (Woolf *et al.*, International PCT Publication No. WO 98/13526; Thompson *et al.*, International PCT Publication No. WO 99/54459; Hartmann *et al.*, USSN 60/101,174, filed on September 21, 1998). All of these references are incorporated by reference herein in their entirety.

In addition, antisense deoxyoligoribonucleotides can be used to target RNA by means of DNA-RNA interactions, thereby activating RNase H, which digests the target RNA in the duplex. Antisense DNA can be expressed via the use of a single stranded DNA intracellular expression vector or equivalents and variations thereof.

RNA interference: RNA interference refers to the process of sequence specific post transcriptional gene silencing in animals mediated by short interfering RNAs (siRNA) (Fire *et al.*, 1998, *Nature*, 391, 806). The corresponding process in plants is commonly referred to as post transcriptional gene silencing or RNA silencing and is also referred to as quelling in fungi. The process of post transcriptional gene silencing is thought to be an evolutionarily conserved cellular defense mechanism used to prevent the expression of foreign genes which is commonly shared by diverse flora and phyla (Fire *et al.*, 1999, *Trends Genet.*, 15, 358). Such protection from foreign gene expression may have evolved in response to the production of double stranded RNAs (dsRNA) derived from viral infection or the random integration of

transposon elements into a host genome via a cellular response that specifically destroys homologous single stranded RNA or viral genomic RNA. The presence of dsRNA in cells triggers the RNAi response through a mechanism that has yet to be fully characterized. This mechanism appears to be different from the interferon response that results from dsRNA mediated activation of protein kinase PKR and 2',5'-oligoadenylate synthetase resulting in non-specific cleavage of mRNA by ribonuclease L.

The presence of long dsRNAs in cells stimulates the activity of a ribonuclease III enzyme referred to as dicer. Dicer is involved in the processing of the dsRNA into short pieces of dsRNA known as short interfering RNAs (siRNA) (Berstein *et al.*, 2001, *Nature*, 409, 363). Short interfering RNAs derived from dicer activity are typically about 21-23 nucleotides in length and comprise about 19 base pair duplexes. Dicer has also been implicated in the excision of 21 and 22 nucleotide small temporal RNAs (stRNA) from precursor RNA of conserved structure that are implicated in translational control (Hutvagner *et al.*, 2001, *Science*, 293, 834). The RNAi response also features an endonuclease complex containing a siRNA, commonly referred to as an RNA-induced silencing complex (RISC), which mediates cleavage of single stranded RNA having sequence homologous to the siRNA. Cleavage of the target RNA takes place in the middle of the region complementary to the guide sequence of the siRNA duplex (Elbashir *et al.*, 2001, *Genes Dev.*, 15, 188).

Short interfering RNA mediated RNAi has been studied in a variety of systems. Fire *et al.*, 1998, *Nature*, 391, 806, were the first to observe RNAi in *C. Elegans*. Wianny and Goetz, 1999, *Nature Cell Biol.*, 2, 70, describes RNAi mediated by dsRNA in mouse embryos. Hammond *et al.*, 2000, *Nature*, 404, 293, describe RNAi in *Drosophila* cells transfected with dsRNA. Elbashir *et al.*, 2001, *Nature*, 411, 494, describe RNAi induced by introduction of duplexes of synthetic 21-nucleotide RNAs in cultured mammalian cells including human embryonic kidney and HeLa cells. Recent work in *Drosophila* embryonic lysates has revealed certain requirements for siRNA length, structure, chemical composition, and sequence that are essential to mediate efficient RNAi activity. These studies have shown that 21 nucleotide siRNA duplexes are most active when containing two nucleotide 3'-overhangs. Furthermore, substitution of one or both siRNA strands with 2'-deoxy or 2'-O-methyl nucleotides abolishes RNAi activity, whereas substitution of 3'-terminal siRNA nucleotides with deoxy nucleotides was shown to be tolerated. Mismatch sequences in the center of the siRNA duplex were also shown to abolish RNAi activity. In addition, these studies also indicate that the position of the cleavage site in the target RNA is defined by the 5'-end of the siRNA guide sequence rather than the 3'-end (Elbashir *et al.*, 2001, *EMBO J.*, 20, 6877). Other studies have indicated that a 5'-phosphate on the target-complementary strand of a siRNA duplex is required for siRNA activity and that ATP is utilized to maintain

the 5'-phosphate moiety on the siRNA (Nykanen *et al.*, 2001, *Cell*, 107, 309), however siRNA molecules lacking a 5'-phosphate are active when introduced exogenously, suggesting that 5'-phosphorylation of siRNA constructs may occur *in vivo*.

Enzymatic Nucleic Acid: Several varieties of naturally-occurring enzymatic RNAs are presently known. In addition, several *in vitro* selection (evolution) strategies (Orgel, 1979, *Proc. R. Soc. London*, B 205, 435) have been used to evolve new nucleic acid catalysts capable of catalyzing cleavage and ligation of phosphodiester linkages (Joyce, 1989, *Gene*, 82, 83-87; Beaudry *et al.*, 1992, *Science* 257, 635-641; Joyce, 1992, *Scientific American* 267, 90-97; Breaker *et al.*, 1994, *TIBTECH* 12, 268; Bartel *et al.*, 1993, *Science* 261:1411-1418; Szostak, 1993, *TIBS* 17, 89-93; Kumar *et al.*, 1995, *FASEB J.*, 9, 1183; Breaker, 1996, *Curr. Op. Biotech.*, 7, 442; Santoro *et al.*, 1997, *Proc. Natl. Acad. Sci.*, 94, 4262; Tang *et al.*, 1997, *RNA* 3, 914; Nakamaye & Eckstein, 1994, *supra*; Long & Uhlenbeck, 1994, *supra*; Ishizaka *et al.*, 1995, *supra*; Vaish *et al.*, 1997, *Biochemistry* 36, 6495; all of these are incorporated by reference herein). Each can catalyze a series of reactions including the hydrolysis of phosphodiester bonds in *trans* (and thus can cleave other RNA molecules) under physiological conditions.

Nucleic acid molecules of this invention can modulate, e.g., down-regulate, Ras protein expression and can be used to treat disease or diagnose disease associated with the levels of Ras, HIV and/or HER2. Enzymatic nucleic acid sequences targeting Ras, HIV and/or HER2 RNA and sequences that can be targeted with nucleic acid molecules of the invention to down-regulate Ras expression are shown in the Tables herein.

The enzymatic nature of an enzymatic nucleic acid molecule allows the concentration of enzymatic nucleic acid molecule necessary to affect a therapeutic treatment to be lower than a nucleic acid molecule lacking enzymatic activity. This reflects the ability of the enzymatic nucleic acid molecule to act enzymatically. Thus, a single enzymatic nucleic acid molecule is able to cleave many molecules of target RNA. In addition, the enzymatic nucleic acid molecule is a highly specific inhibitor, with the specificity of inhibition depending not only on the base-pairing mechanism of binding to the target RNA, but also on the mechanism of target RNA cleavage. Single mismatches, or base-substitutions, near the site of cleavage can be chosen to completely eliminate catalytic activity of a enzymatic nucleic acid molecule.

Nucleic acid molecules having an endonuclease enzymatic activity are able to repeatedly cleave other separate RNA molecules in a nucleotide base sequence-specific manner. With proper design and construction, such enzymatic nucleic acid molecules can be targeted to virtually any RNA transcript, and achieve efficient cleavage *in vitro* (Zaug *et al.*,

324, *Nature* 429 1986; Uhlenbeck, 1987 *Nature* 328, 596; Kim *et al.*, 84 *Proc. Natl. Acad. Sci. USA* 8788, 1987; Dreyfus, 1988, *Einstein Quart. J. Bio. Med.*, 6, 92; Haseloff and Gerlach, 334 *Nature* 585, 1988; Cech, 260 *JAMA* 3030, 1988; and Jefferies *et al.*, 17 *Nucleic Acids Research* 1371, 1989; Santoro *et al.*, 1997 *supra*).

5 Because of their sequence specificity, *trans*-cleaving enzymatic nucleic acid molecules can be used as therapeutic agents for human disease (Usman & McSwiggen, 1995 *Ann. Rep. Med. Chem.* 30, 285-294; Christoffersen and Marr, 1995 *J. Med. Chem.* 38, 2023-2037). Enzymatic nucleic acid molecules can be designed to cleave specific RNA targets within the background of cellular RNA. Such a cleavage event renders the RNA non-functional and
10 abrogates protein expression from that RNA. In this manner, synthesis of a protein associated with a disease state can be selectively inhibited (Warashina *et al.*, 1999, *Chemistry and Biology*, 6, 237-250).

Enzymatic nucleic acid molecules of the invention that are allosterically regulated ("allozymes") can be used to modulate, including down-regulate, Ras, HIV and/or HER2
15 expression. These allosteric enzymatic nucleic acids or allozymes (see for example George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, International PCT publication No. WO 99/29842) are designed to
20 respond to a signaling agent, for example, mutant Ras, HIV and/or HER2 protein, wild-type Ras, HIV and/or HER2 protein, mutant Ras, HIV and/or HER2 RNA, wild-type Ras, HIV and/or HER2 RNA, other proteins and/or RNAs involved in Ras, HIV and/or HER2 activity, compounds, metals, polymers, molecules and/or drugs that are targeted to Ras, HIV and/or HER2 expressing cells etc., which, in turn, modulate the activity of the enzymatic nucleic
25 acid molecule. In response to interaction with a predetermined signaling agent, the activity of the allosteric enzymatic nucleic acid molecule is activated or inhibited such that the expression of a particular target is selectively regulated, including down-regulated. The target can comprise wild-type Ras, HIV and/or HER2, mutant Ras, HIV and/or HER2, a component of Ras, HIV and/or HER2, and/or a predetermined cellular component that modulates Ras,
30 HIV and/or HER2 activity. For example, allosteric enzymatic nucleic acid molecules that are activated by interaction with a RNA encoding Ras, HIV and/or HER2 protein can be used as therapeutic agents *in vivo*. The presence of RNA encoding the Ras, HIV and/or HER2 protein activates the allosteric enzymatic nucleic acid molecule that subsequently cleaves the RNA encoding Ras, HIV and/or HER2 protein, resulting in the inhibition of Ras, HIV and/or

HER2 protein expression. In this manner, cells that express the Ras, HIV and/or HER2 protein are selectively targeted.

In another non-limiting example, an allozyme can be activated by a Ras, HIV and/or HER2 protein, peptide, or mutant polypeptide that causes the allozyme to inhibit the expression of Ras, HIV and/or HER2 gene, by, for example, cleaving RNA encoded by Ras, HIV and/or HER2 gene. In this non-limiting example, the allozyme acts as a decoy to inhibit the function of Ras, HIV and/or HER2 and also inhibit the expression of Ras, HIV and/or HER2 once activated by the Ras, HIV and/or HER2 protein.

Target sites

Targets for useful enzymatic nucleic acid molecules and antisense nucleic acids can be determined as disclosed in Draper *et al.*, WO 93/23569; Sullivan *et al.*, WO 93/23057; Thompson *et al.*, WO 94/02595; Draper *et al.*, WO 95/04818; McSwiggen *et al.*, US Patent No. 5,525,468, and hereby incorporated by reference herein in totality. Other examples include the following PCT applications, which concern inactivation of expression of disease-related genes: WO 95/23225, WO 95/13380, WO 94/02595, incorporated by reference herein. Rather than repeat the guidance provided in those documents here, below are provided specific non-limiting examples of such methods. Enzymatic nucleic acid molecules to such targets are designed as described in the above applications and synthesized to be tested *in vitro* and *in vivo*, as also described. The sequences of human K-Ras, H-Ras, HIV-1 and HER2 RNAs were screened for optimal enzymatic nucleic acid target sites using a computer-folding algorithm. Nucleic acid molecule binding/cleavage sites were identified. These sites are shown in the Tables (all sequences are 5' to 3' in the tables). The nucleotide base position is noted in the Tables as that site to be cleaved by the designated type of enzymatic nucleic acid molecule. Human sequences can be screened and enzymatic nucleic acid molecule and/or antisense thereafter designed, as discussed in Stinchcomb *et al.*, WO 95/23225. In addition, mouse targeted nucleic acid molecules can be used to test efficacy of action of the enzymatic nucleic acid molecule, siRNA and/or antisense prior to testing in humans.

In addition, enzymatic nucleic acid, siRNA, and antisense nucleic acid molecule binding/cleavage sites were identified. The nucleic acid molecules are individually analyzed by computer folding (Jaeger *et al.*, 1989 *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether the sequences fold into the appropriate secondary structure. Those nucleic acid molecules with unfavorable intramolecular interactions, such as between, for example the binding arms and the catalytic core of an enzymatic nucleic acid, are eliminated from consideration. Varying binding arm lengths can be chosen to optimize activity.

Antisense, hammerhead, DNAzyme, NCH, amberzyme, zinzyme or G-Cleaver enzymatic nucleic acid molecule, siRNA, and antisense nucleic acid binding/cleavage sites were identified and were designed to anneal to various sites in the RNA target. The enzymatic nucleic acid binding arms or siRNA and antisense nucleic acid sequences are complementary to the target site sequences described above. The nucleic acid molecules are chemically synthesized. The method of synthesis used follows the procedure for normal DNA/RNA synthesis as described below and in Usman *et al.*, 1987 *J. Am. Chem. Soc.*, 109, 7845; Scaringe *et al.*, 1990 *Nucleic Acids Res.*, 18, 5433; and Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677-2684; Caruthers *et al.*, 1992, *Methods in Enzymology* 211, 3-19.

10 Synthesis of Nucleic acid Molecules

Synthesis of nucleic acids greater than 100 nucleotides in length can be difficult using automated methods, and the therapeutic cost of such molecules can be prohibitive. In this invention, small nucleic acid motifs ("small" refers to nucleic acid motifs less than about 100 nucleotides in length, preferably less than about 80 nucleotides in length, and more preferably less than about 50 nucleotides in length; *e.g.*, DNAzymes) are preferably used for exogenous delivery. The simple structure of these molecules increases the ability of the nucleic acid to invade targeted regions of RNA structure. Exemplary molecules of the instant invention are chemically synthesized as described herein, and others can similarly be synthesized.

Oligonucleotides (*e.g.*, DNAzymes, antisense) are synthesized using protocols known in the art as described in Caruthers *et al.*, 1992, *Methods in Enzymology* 211, 3-19, Thompson *et al.*, International PCT Publication No. WO 99/54459, Wincott *et al.*, 1995, *Nucleic Acids Res.* 23, 2677-2684, Wincott *et al.*, 1997, *Methods Mol. Bio.*, 74, 59, Brennan *et al.*, 1998, *Biotechnol Bioeng.*, 61, 33-45, and Brennan, US patent No. 6,001,311. All of these references are incorporated herein by reference. The synthesis of oligonucleotides makes use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. In a non-limiting example, small scale syntheses are conducted on a 394 Applied Biosystems, Inc. synthesizer using a 0.2 μ mol scale protocol with a 2.5 min coupling step for 2'-O-methylated nucleotides and a 45 sec coupling step for 2'-deoxy nucleotides. **Table I** outlines the amounts and the contact times of the reagents used in the synthesis cycle. Alternatively, syntheses at the 0.2 μ mol scale can be performed on a 96-well plate synthesizer, such as the instrument produced by Protogene (Palo Alto, CA) with minimal modification to the cycle. A 33-fold excess (60 μ L of 0.11 M = 6.6 μ mol) of 2'-O-methyl phosphoramidite and a 105-fold excess of S-ethyl tetrazole (60 μ L of 0.25 M = 15 μ mol) can be used in each coupling cycle of 2'-O-methyl residues relative to polymer-

- bound 5'-hydroxyl. A 22-fold excess (40 μ L of 0.11 M = 4.4 μ mol) of deoxy phosphoramidite and a 70-fold excess of S-ethyl tetrazole (40 μ L of 0.25 M = 10 μ mol) can be used in each coupling cycle of deoxy residues relative to polymer-bound 5'-hydroxyl. Average coupling yields on the 394 Applied Biosystems, Inc. synthesizer, determined by colorimetric quantitation of the trityl fractions, are typically 97.5-99%. Other oligonucleotide synthesis reagents for the 394 Applied Biosystems, Inc. synthesizer include; detritylation solution is 3% TCA in methylene chloride (ABI); capping is performed with 16% *N*-methyl imidazole in THF (ABI) and 10% acetic anhydride/10% 2,6-lutidine in THF (ABI); and oxidation solution is 16.9 mM I₂, 49 mM pyridine, 9% water in THF (PERSEPTIVE™).
- 10 Burdick & Jackson Synthesis Grade acetonitrile is used directly from the reagent bottle. S-Ethyltetrazole solution (0.25 M in acetonitrile) is made up from the solid obtained from American International Chemical, Inc. Alternately, for the introduction of phosphorothioate linkages, Beaucage reagent (3H-1,2-Benzodithiol-3-one 1,1-dioxide, 0.05 M in acetonitrile) is used.
- 15 Deprotection of the DNazymes is performed as follows: the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 40% aq. methylamine (1 mL) at 65 °C for 10 min. After cooling to -20 °C, the supernatant is removed from the polymer support. The support is washed three times with 1.0 mL of EtOH:MeCN:H₂O/3:1:1, vortexed and the supernatant is then added to the first supernatant.
- 20 The combined supernatants, containing the oligoribonucleotide, are dried to a white powder.

- The method of synthesis used for RNA and chemically modified RNA or DNA, including certain enzymatic nucleic acid molecules and siRNA molecules, follows the procedure as described in Usman *et al.*, 1987, *J. Am. Chem. Soc.*, 109, 7845; Scaringe *et al.*, 1990, *Nucleic Acids Res.*, 18, 5433; and Wincott *et al.*, 1995, *Nucleic Acids Res.* 23, 2677-2684 Wincott *et al.*, 1997, *Methods Mol. Bio.*, 74, 59, and makes use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. In a non-limiting example, small scale syntheses are conducted on a 394 Applied Biosystems, Inc. synthesizer using a 0.2 μ mol scale protocol with a 7.5 min coupling step for alkylsilyl protected nucleotides and a 2.5 min coupling step for 2'-O-methylated nucleotides. Table I outlines the amounts and the contact times of the reagents used in the synthesis cycle. Alternatively, syntheses at the 0.2 μ mol scale can be done on a 96-well plate synthesizer, such as the instrument produced by Protogene (Palo Alto, CA) with minimal modification to the cycle. A 33-fold excess (60 μ L of 0.11 M = 6.6 μ mol) of 2'-O-methyl phosphoramidite and a 75-fold excess of S-ethyl tetrazole (60 μ L of 0.25 M = 15 μ mol) can be used in each coupling cycle of 2'-O-methyl residues relative to polymer-bound 5'-
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hydroxyl. A 66-fold excess (120 μL of 0.11 M = 13.2 μmol) of alkylsilyl (ribo) protected phosphoramidite and a 150-fold excess of S-ethyl tetrazole (120 μL of 0.25 M = 30 μmol) can be used in each coupling cycle of ribo residues relative to polymer-bound 5'-hydroxyl. Average coupling yields on the 394 Applied Biosystems, Inc. synthesizer, determined by colorimetric quantitation of the trityl fractions, are typically 97.5-99%. Other oligonucleotide synthesis reagents for the 394 Applied Biosystems, Inc. synthesizer include; detritylation solution is 3% TCA in methylene chloride (ABI); capping is performed with 16% *N*-methyl imidazole in THF (ABI) and 10% acetic anhydride/10% 2,6-lutidine in THF (ABI); oxidation solution is 16.9 mM I_2 , 49 mM pyridine, 9% water in THF (PERSEPTIVETM). Burdick & Jackson Synthesis Grade acetonitrile is used directly from the reagent bottle. S-Ethyltetrazole solution (0.25 M in acetonitrile) is made up from the solid obtained from American International Chemical, Inc. Alternately, for the introduction of phosphorothioate linkages, Beaucage reagent (3H-1,2-Benzodithiol-3-one 1,1-dioxide 0.05 M in acetonitrile) is used.

Deprotection of the RNA is performed using either a two-pot or one-pot protocol. For the two-pot protocol, the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 40% aq. methylamine (1 mL) at 65 °C for 10 min. After cooling to -20 °C, the supernatant is removed from the polymer support. The support is washed three times with 1.0 mL of EtOH:MeCN:H₂O/3:1:1, vortexed and the supernatant is then added to the first supernatant. The combined supernatants, containing the oligoribonucleotide, are dried to a white powder. The base deprotected oligoribonucleotide is resuspended in anhydrous TEA/HF/NMP solution (300 μL of a solution of 1.5 mL *N*-methylpyrrolidinone, 750 μL TEA and 1 mL TEA•3HF to provide a 1.4 M HF concentration) and heated to 65 °C. After 1.5 h, the oligomer is quenched with 1.5 M NH_4HCO_3 .

Alternatively, for the one-pot protocol, the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 33% ethanolic methylamine/DMSO: 1/1 (0.8 mL) at 65 °C for 15 min. The vial is brought to r.t. TEA•3HF (0.1 mL) is added and the vial is heated at 65 °C for 15 min. The sample is cooled at -20 °C and then quenched with 1.5 M NH_4HCO_3 .

For purification of the trityl-on oligomers, the quenched NH_4HCO_3 solution is loaded onto a C-18 containing cartridge that had been prewashed with acetonitrile followed by 50 mM TEAA. After washing the loaded cartridge with water, the RNA is detritylated with 0.5% TFA for 13 min. The cartridge is then washed again with water, salt exchanged with 1 M NaCl and washed with water again. The oligonucleotide is then eluted with 30% acetonitrile.

Inactive nucleic acid molecules or binding attenuated control (BAC) oligonucleotides can be synthesized by substituting one or more nucleotides in the nucleic acid molecule to inactivate the molecule and such molecules can serve as a negative control.

The average stepwise coupling yields are typically >98% (Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677-2684). Those of ordinary skill in the art will recognize that the scale of synthesis can be adapted to be larger or smaller than the example described above including but not limited to 96 well format, all that is important is the ratio of chemicals used in the reaction.

Alternatively, the nucleic acid molecules of the present invention can be synthesized separately and joined together post-synthetically, for example by ligation (Moore *et al.*, 1992, *Science* 256, 9923; Draper *et al.*, International PCT publication No. WO 93/23569; Shabarova *et al.*, 1991, *Nucleic Acids Research* 19, 4247; Bellon *et al.*, 1997, *Nucleosides & Nucleotides*, 16, 951; Bellon *et al.*, 1997, *Bioconjugate Chem.* 8, 204).

The nucleic acid molecules of the present invention can be modified extensively to enhance stability by modification with nuclease resistant groups, for example, 2'-amino, 2'-C-allyl, 2'-fluoro, 2'-O-methyl, 2'-H (for a review see Usman and Cedergren, 1992, *TIBS* 17, 34; Usman *et al.*, 1994, *Nucleic Acids Symp. Ser.* 31, 163). Enzymatic nucleic acid molecules are purified by gel electrophoresis using known methods or are purified by high pressure liquid chromatography (HPLC; See Wincott *et al.*, *Supra*, the totality of which is hereby incorporated herein by reference) and are re-suspended in water.

The sequences of the nucleic acid molecules, including enzymatic nucleic acid molecules and antisense, that are chemically synthesized, are shown in the Tables herein. These sequences are representative only of many more such sequences where the enzymatic portion of the enzymatic nucleic acid molecule (all but the binding arms) is modified to affect activity. For example, the enzymatic nucleic acid sequences listed in the Tables can be formed of deoxyribonucleotides or other nucleotides or non-nucleotides. Such enzymatic nucleic acid molecules with enzymatic activity are equivalent to the enzymatic nucleic acid molecules described specifically in the Tables.

Optimizing Activity of the Nucleic Acid Molecule of the Invention.

Chemically synthesizing nucleic acid molecules with modifications (base, sugar and/or phosphate) that prevent their degradation by serum ribonucleases can increase their potency (see *e.g.*, Eckstein *et al.*, International Publication No. WO 92/07065; Perrault *et al.*, 1990 *Nature* 344, 565; Pieken *et al.*, 1991, *Science* 253, 314; Usman and Cedergren, 1992, *Trends*

in *Biochem. Sci.* 17, 334; Usman *et al.*, International Publication No. WO 93/15187; and Rossi *et al.*, International Publication No. WO 91/03162; Sproat, US Patent No. 5,334,711; and Burgin *et al.*, *supra*, all of which are hereby incorporated by reference in their entirety). All of the above references describe various chemical modifications that can be made to the base, phosphate and/or sugar moieties of the nucleic acid molecules described herein. Modifications which enhance their efficacy in cells, and removal of bases from nucleic acid molecules to shorten oligonucleotide synthesis times and reduce chemical requirements are desired.

There are several examples of sugar, base and phosphate modifications that can be introduced into nucleic acid molecules with significant enhancement in their nuclease stability and efficacy. For example, oligonucleotides can be modified to enhance stability and/or enhance biological activity by modification with nuclease resistant groups, for example, 2'-amino, 2'-C-allyl, 2'-fluoro, 2'-O-methyl, 2'-H, nucleotide base modifications (for a review see Usman and Cedergren, 1992, *TIBS*, 17, 34; Usman *et al.*, 1994, *Nucleic Acids Symp. Ser.* 31, 163; Burgin *et al.*, 1996, *Biochemistry*, 35, 14090). Sugar modification of nucleic acid molecules are also known to increase efficacy (see Eckstein *et al.*, International Publication PCT No. WO 92/07065; Perrault *et al.*, *Nature*, 1990, 344, 565-568; Pieken *et al.*, *Science*, 1991, 253, 314-317; Usman and Cedergren, *Trends in Biochem. Sci.*, 1992, 17, 334-339; Usman *et al.*, International Publication PCT No. WO 93/15187; Sproat, US Patent No. 5,334,711 and Beigelman *et al.*, 1995, *J. Biol. Chem.*, 270, 25702; Beigelman *et al.*, International PCT publication No. WO 97/26270; Beigelman *et al.*, US Patent No. 5,716,824; Usman *et al.*, US patent No. 5,627,053; Woolf *et al.*, International PCT Publication No. WO 98/13526; Thompson *et al.*, USSN 60/082,404 which was filed on April 20, 1998; Karpeisky *et al.*, 1998, *Tetrahedron Lett.*, 39, 1131; Earnshaw and Gait, 1998, *Biopolymers (Nucleic acid Sciences)*, 48, 39-55; Verma and Eckstein, 1998, *Annu. Rev. Biochem.*, 67, 99-134; and Burlina *et al.*, 1997, *Bioorg. Med. Chem.*, 5, 1999-2010; all of the references are hereby incorporated in their totality by reference herein). The publications describe general methods and strategies to determine the location of incorporation of sugar, base and/or phosphate modifications and the like into enzymatic nucleic acid molecules without inhibiting catalysis. Similar modifications can be used as described herein to modify the nucleic acid molecules of the instant invention.

While chemical modification of oligonucleotide internucleotide linkages with phosphorothioate, phosphorothioate, and/or 5'-methylphosphonate linkages improves stability, excessive modifications can cause some toxicity. Therefore, when designing nucleic acid molecules, the amount of these internucleotide linkages should be minimized. The

reduction in the concentration of these linkages can lower toxicity, resulting in increased efficacy and higher specificity of the therapeutic nucleic acid molecules.

Nucleic acid molecules having chemical modifications that maintain or enhance activity are provided. Such nucleic acid molecules are also generally more resistant to nucleases than unmodified nucleic acid molecules. Thus, the *in vitro* and/or *in vivo* activity should not be significantly lowered. Therapeutic nucleic acid molecules delivered exogenously are optimally stable within cells until translation of the target RNA has been inhibited long enough to reduce the levels of the undesirable protein. This period of time varies between hours to days, depending upon the disease state. Nucleic acid molecules are preferably resistant to nucleases in order to function as effective intracellular therapeutic agents. Improvements in the chemical synthesis of RNA and DNA (Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677; Caruthers *et al.*, 1992, *Methods in Enzymology* 211,3-19 (incorporated by reference herein)) have expanded the ability to modify nucleic acid molecules by introducing nucleotide modifications to enhance their nuclease stability as described above.

In one embodiment, nucleic acid molecules of the invention include one or more G-clamp nucleotides. A G-clamp nucleotide is a modified cytosine analog wherein modifications result in the ability to hydrogen bond both Watson-Crick and Hoogsteen faces of a complementary guanine within a duplex, see for example Lin and Matteucci, 1998, *J. Am. Chem. Soc.*, 120, 8531-8532. A single G-clamp analog substitution within an oligonucleotide can result in substantially enhanced helical thermal stability and mismatch discrimination when hybridized to complementary oligonucleotides. The inclusion of such nucleotides in nucleic acid molecules of the invention can enable both enhanced affinity and specificity to nucleic acid targets.

In another embodiment, the invention features conjugates and/or complexes of nucleic acid molecules targeting Ras genes such as K-Ras, H-Ras, and/or N-Ras. Compositions and conjugates are used to facilitate delivery of molecules into a biological system, such as cells. The conjugates provided by the instant invention can impart therapeutic activity by transferring therapeutic compounds across cellular membranes, altering the pharmacokinetics, and/or modulating the localization of nucleic acid molecules of the invention. The present invention encompasses the design and synthesis of novel agents for the delivery of molecules, including but not limited to, small molecules, lipids, phospholipids, nucleosides, nucleotides, nucleic acids, antibodies, toxins, negatively charged polymers and other polymers, for example proteins, peptides, hormones, carbohydrates, polyethylene glycols, or polyamines, across cellular membranes. In general, the transporters described are designed to be used

either individually or as part of a multi-component system, with or without degradable linkers. These compounds are expected to improve delivery and/or localization of nucleic acid molecules of the invention into a number of cell types originating from different tissues, in the presence or absence of serum (see Sullenger and Cech, US 5,854,038). Conjugates of the molecules described herein can be attached to biologically active molecules via linkers that are biodegradable, such as biodegradable nucleic acid linker molecules.

The term "biodegradable nucleic acid linker molecule" as used herein, refers to a nucleic acid molecule that is designed as a biodegradable linker to connect one molecule to another molecule, for example, a biologically active molecule. The stability of the biodegradable nucleic acid linker molecule can be modulated by using various combinations of ribonucleotides, deoxyribonucleotides, and chemically modified nucleotides, for example 2'-O-methyl, 2'-fluoro, 2'-amino, 2'-O-amino, 2'-C-allyl, 2'-O-allyl, and other 2'-modified or base modified nucleotides. The biodegradable nucleic acid linker molecule can be a dimer, trimer, tetramer or longer nucleic acid molecule, for example, an oligonucleotide of about 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length, or can comprise a single nucleotide with a phosphorus based linkage, for example, a phosphoramidate or phosphodiester linkage. The biodegradable nucleic acid linker molecule can also comprise nucleic acid backbone, nucleic acid sugar, or nucleic acid base modifications.

The term "biodegradable" as used herein, refers to degradation in a biological system, for example, enzymatic degradation or chemical degradation.

The term "biologically active molecule" as used herein, refers to compounds or molecules that are capable of eliciting or modifying a biological response in a system. Non-limiting examples of biologically active molecules contemplated by the instant invention include therapeutically active molecules such as antibodies, hormones, antivirals, peptides, proteins, chemotherapeutics, small molecules, vitamins, co-factors, nucleosides, nucleotides, oligonucleotides, enzymatic nucleic acids, antisense nucleic acids, triplex forming oligonucleotides, 2,5-A chimeras, siRNA, dsRNA, allozymes, aptamers, decoys and analogs thereof. Biologically active molecules of the invention also include molecules capable of modulating the pharmacokinetics and/or pharmacodynamics of other biologically active molecules, for example lipids and polymers such as polyamines, polyamides, polyethylene glycol and other polyethers.

The term "phospholipid" as used herein, refers to a hydrophobic molecule comprising at least one phosphorus group. For example, a phospholipid can comprise a phosphorus

containing group and saturated or unsaturated alkyl group, optionally substituted with OH, COOH, oxo, amine, or substituted or unsubstituted aryl groups.

Use of the nucleic acid-based molecules of the invention can lead to better treatment of the disease progression by affording the possibility of combination therapies (*e.g.*, multiple antisense or enzymatic nucleic acid molecules targeted to different genes, nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of molecules (including different motifs) and/or other chemical or biological molecules). The treatment of subjects with nucleic acid molecules can also include combinations of different types of nucleic acid molecules.

In the case that down-regulation of the target is desired, therapeutic nucleic acid molecules (*e.g.*, DNAzymes) delivered exogenously are optimally stable within cells until translation of the target RNA has been inhibited long enough to reduce the levels of the targeted protein. This period of time varies between hours to days depending upon the disease state. These nucleic acid molecules should be resistant to nucleases in order to function as effective intracellular therapeutic agents. Improvements in the chemical synthesis of nucleic acid molecules described in the instant invention and others known in the art have expanded the ability to modify nucleic acid molecules by introducing nucleotide modifications to enhance their nuclease stability as described above.

In another embodiment, nucleic acid catalysts having chemical modifications that maintain or enhance enzymatic activity are provided. Such nucleic acids are also generally more resistant to nucleases than unmodified nucleic acid. Thus, the *in vitro* and/or *in vivo* the activity of the nucleic acid should not be significantly lowered. As exemplified herein, such enzymatic nucleic acids are useful for *in vitro* and/or *in vivo* techniques even if activity over all is reduced 10 fold (Burgin *et al.*, 1996, *Biochemistry*, 35, 14090). Such enzymatic nucleic acids herein are said to "maintain" the enzymatic activity of an all RNA ribozyme or all DNA DNAzyme.

In another aspect the nucleic acid molecules comprise a 5' and/or a 3'- cap structure.

By "cap structure" is meant chemical modifications, which have been incorporated at either terminus of the oligonucleotide (see, for example, Wincott *et al.*, WO 97/26270, incorporated by reference herein). These terminal modifications protect the nucleic acid molecule from exonuclease degradation, and can help in delivery and/or localization within a cell. The cap can be present at the 5'-terminus (5'-cap) or at the 3'-terminus (3'-cap) or can be present on both termini. In non-limiting examples, the 5'-cap includes inverted abasic

residue (moiety), 4',5'-methylene nucleotide; 1-(beta-D-erythrofuransyl) nucleotide, 4'-thio nucleotide, carbocyclic nucleotide; 1,5-anhydrohexitol nucleotide; L-nucleotides; alpha-nucleotides; modified base nucleotide; phosphorodithioate linkage; *threo*-pentofuransyl nucleotide; acyclic 3',4'-seco nucleotide; acyclic 3,4-dihydroxybutyl nucleotide; acyclic 3,5-dihydroxypentyl nucleotide, 3'-3'-inverted nucleotide moiety; 3'-3'-inverted abasic moiety; 3'-2'-inverted nucleotide moiety; 3'-2'-inverted abasic moiety; 1,4-butanediol phosphate; 3'-phosphoramidate; hexylphosphate; aminohexyl phosphate; 3'-phosphate; 3'-phosphorothioate; phosphorodithioate; or bridging or non-bridging methylphosphonate moiety (for more details see Wincott *et al.*, International PCT publication No. WO 97/26270, incorporated by reference herein).

In another embodiment, the 3'-cap includes, for example 4',5'-methylene nucleotide; 1-(beta-D-erythrofuransyl) nucleotide; 4'-thio nucleotide, carbocyclic nucleotide; 5'-amino-alkyl phosphate; 1,3-diamino-2-propyl phosphate, 3-aminopropyl phosphate; 6-aminohexyl phosphate; 1,2-aminododecyl phosphate; hydroxypropyl phosphate; 1,5-anhydrohexitol nucleotide; L-nucleotide; alpha-nucleotide; modified base nucleotide; phosphorodithioate; *threo*-pentofuransyl nucleotide; acyclic 3',4'-seco nucleotide; 3,4-dihydroxybutyl nucleotide; 3,5-dihydroxypentyl nucleotide, 5'-5'-inverted nucleotide moiety; 5'-5'-inverted abasic moiety; 5'-phosphoramidate; 5'-phosphorothioate; 1,4-butanediol phosphate; 5'-amino; bridging and/or non-bridging 5'-phosphoramidate, phosphorothioate and/or phosphorodithioate, bridging or non bridging methylphosphonate and 5'-mercapto moieties (for more details see Beaucage and Iyer, 1993, *Tetrahedron* 49, 1925; incorporated by reference herein).

By the term "non-nucleotide" is meant any group or compound which can be incorporated into a nucleic acid chain in the place of one or more nucleotide units, including either sugar and/or phosphate substitutions, and allows the remaining bases to exhibit their enzymatic activity. The group or compound is abasic in that it does not contain a commonly recognized nucleotide base, such as adenosine, guanine, cytosine, uracil or thymine.

The term "alkyl" as used herein refers to a saturated aliphatic hydrocarbon, including straight-chain, branched-chain "isoalkyl", and cyclic alkyl groups. The term "alkyl" also comprises alkoxy, alkyl-thio, alkyl-thio-alkyl, alkoxyalkyl, alkylamino, alkenyl, alkynyl, alkoxy, cycloalkenyl, cycloalkyl, cycloalkylalkyl, heterocycloalkyl, heteroaryl, C1-C6 hydrocarbyl, aryl or substituted aryl groups. Preferably, the alkyl group has 1 to 12 carbons. More preferably it is a lower alkyl of from about 1 to 7 carbons, more preferably about 1 to 4 carbons. The alkyl group can be substituted or unsubstituted. When substituted the substituted group(s) preferably comprise hydroxy, oxy, thio, amino, nitro, cyano, alkoxy,

alkyl-thio, alkyl-thio-alkyl, alkoxyalkyl, alkylamino, silyl, alkenyl, alkynyl, alkoxy, cycloalkenyl, cycloalkyl, cycloalkylalkyl, heterocycloalkyl, heteroaryl, C1-C6 hydrocarbyl, aryl or substituted aryl groups. The term "alkyl" also includes alkenyl groups containing at least one carbon-carbon double bond, including straight-chain, branched-chain, and cyclic groups. Preferably, the alkenyl group has about 2 to 12 carbons. More preferably it is a lower alkenyl of from about 2 to 7 carbons, more preferably about 2 to 4 carbons. The alkenyl group can be substituted or unsubstituted. When substituted the substituted group(s) preferably comprise hydroxy, oxy, thio, amino, nitro, cyano, alkoxy, alkyl-thio, alkyl-thio-alkyl, alkoxyalkyl, alkylamino, silyl, alkenyl, alkynyl, alkoxy, cycloalkenyl, cycloalkyl, cycloalkylalkyl, heterocycloalkyl, heteroaryl, C1-C6 hydrocarbyl, aryl or substituted aryl groups.

The term "alkyl" also includes alkynyl groups containing at least one carbon-carbon triple bond, including straight-chain, branched-chain, and cyclic groups. Preferably, the alkynyl group has about 2 to 12 carbons. More preferably it is a lower alkynyl of from about 2 to 7 carbons, more preferably about 2 to 4 carbons. The alkynyl group can be substituted or unsubstituted. When substituted the substituted group(s) preferably comprise hydroxy, oxy, thio, amino, nitro, cyano, alkoxy, alkyl-thio, alkyl-thio-alkyl, alkoxyalkyl, alkylamino, silyl, alkenyl, alkynyl, alkoxy, cycloalkenyl, cycloalkyl, cycloalkylalkyl, heterocycloalkyl, heteroaryl, C1-C6 hydrocarbyl, aryl or substituted aryl groups. Alkyl groups or moieties of the invention can also include aryl, alkylaryl, carbocyclic aryl, heterocyclic aryl, amide and ester groups. The preferred substituent(s) of aryl groups are halogen, trihalomethyl, hydroxyl, SH, OH, cyano, alkoxy, alkyl, alkenyl, alkynyl, and amino groups. An "alkylaryl" group refers to an alkyl group (as described above) covalently joined to an aryl group (as described above). Carbocyclic aryl groups are groups wherein the ring atoms on the aromatic ring are all carbon atoms. The carbon atoms are optionally substituted. Heterocyclic aryl groups are groups having from about 1 to 3 heteroatoms as ring atoms in the aromatic ring and the remainder of the ring atoms are carbon atoms. Suitable heteroatoms include oxygen, sulfur, and nitrogen, and include furanyl, thienyl, pyridyl, pyrrolyl, N-lower alkyl pyrrolo, pyrimidyl, pyrazinyl, imidazolyl and the like, all optionally substituted. An "amide" refers to an -C(O)-NH-R, where R is either alkyl, aryl, alkylaryl or hydrogen. An "ester" refers to an -C(O)-OR', where R is either alkyl, aryl, alkylaryl or hydrogen.

The term "alkoxyalkyl" as used herein refers to an alkyl-O-alkyl ether, for example, methoxyethyl or ethoxymethyl.

The term "alkyl-thio-alkyl" as used herein refers to an alkyl-S-alkyl thioether, for example, methylthiomethyl or methylthioethyl.

5 The term "amino" as used herein refers to a nitrogen containing group as is known in the art derived from ammonia by the replacement of one or more hydrogen radicals by organic radicals. For example, the terms "aminoacyl" and "aminoalkyl" refer to specific N-substituted organic radicals with acyl and alkyl substituent groups respectively.

The term "amination" as used herein refers to a process in which an amino group or substituted amine is introduced into an organic molecule.

10 The term "exocyclic amine protecting moiety" as used herein refers to a nucleobase amino protecting group compatible with oligonucleotide synthesis, for example, an acyl or amide group.

The term "alkenyl" as used herein refers to a straight or branched hydrocarbon of a designed number of carbon atoms containing at least one carbon-carbon double bond. Examples of "alkenyl" include vinyl, allyl, and 2-methyl-3-heptene.

15 The term "alkoxy" as used herein refers to an alkyl group of indicated number of carbon atoms attached to the parent molecular moiety through an oxygen bridge. Examples of alkoxy groups include, for example, methoxy, ethoxy, propoxy and isopropoxy.

20 The term "alkynyl" as used herein refers to a straight or branched hydrocarbon of a designed number of carbon atoms containing at least one carbon-carbon triple bond. Examples of "alkynyl" include propargyl, propyne, and 3-hexyne.

25 The term "aryl" as used herein refers to an aromatic hydrocarbon ring system containing at least one aromatic ring. The aromatic ring can optionally be fused or otherwise attached to other aromatic hydrocarbon rings or non-aromatic hydrocarbon rings. Examples of aryl groups include, for example, phenyl, naphthyl, 1,2,3,4-tetrahydronaphthalene and biphenyl. Preferred examples of aryl groups include phenyl and naphthyl.

The term "cycloalkenyl" as used herein refers to a C3-C8 cyclic hydrocarbon containing at least one carbon-carbon double bond. Examples of cycloalkenyl include cyclopropenyl, cyclobutenyl, cyclopentenyl, cyclopentadiene, cyclohexenyl, 1,3-cyclohexadiene, cycloheptenyl, cycloheptatrienyl, and cyclooctenyl.

The term "cycloalkyl" as used herein refers to a C3-C8 cyclic hydrocarbon. Examples of cycloalkyl include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl and cyclooctyl.

5 The term "cycloalkylalkyl," as used herein, refers to a C3-C7 cycloalkyl group attached to the parent molecular moiety through an alkyl group, as defined above. Examples of cycloalkylalkyl groups include cyclopropylmethyl and cyclopentylethyl.

The terms "halogen" or "halo" as used herein refers to indicate fluorine, chlorine, bromine, and iodine.

10 The term "heterocycloalkyl," as used herein refers to a non-aromatic ring system containing at least one heteroatom selected from nitrogen, oxygen, and sulfur. The heterocycloalkyl ring can be optionally fused to or otherwise attached to other heterocycloalkyl rings and/or non-aromatic hydrocarbon rings. Preferred heterocycloalkyl groups have from 3 to 7 members. Examples of heterocycloalkyl groups include, for example, piperazine, morpholine, piperidine, tetrahydrofuran, pyrrolidine, and pyrazole.
15 Preferred heterocycloalkyl groups include piperidinyl, piperazinyl, morpholinyl, and pyrrolidinyl.

The term "heteroaryl" as used herein refers to an aromatic ring system containing at least one heteroatom selected from nitrogen, oxygen, and sulfur. The heteroaryl ring can be fused or otherwise attached to one or more heteroaryl rings, aromatic or non-aromatic
20 hydrocarbon rings or heterocycloalkyl rings. Examples of heteroaryl groups include, for example, pyridine, furan, thiophene, 5,6,7,8-tetrahydroisoquinoline and pyrimidine. Preferred examples of heteroaryl groups include thienyl, benzothienyl, pyridyl, quinolyl, pyrazinyl, pyrimidyl, imidazolyl, benzimidazolyl, furanyl, benzofuranyl, thiazolyl, benzothiazolyl, isoxazolyl, oxadiazolyl, isothiazolyl, benzisothiazolyl, triazolyl, tetrazolyl, pyrrolyl, indolyl,
25 pyrazolyl, and benzopyrazolyl.

The term "C1-C6 hydrocarbyl" as used herein refers to straight, branched, or cyclic alkyl groups having 1-6 carbon atoms, optionally containing one or more carbon-carbon double or triple bonds. Examples of hydrocarbyl groups include, for example, methyl, ethyl, propyl, isopropyl, n-butyl, sec-butyl, tert-butyl, pentyl, 2-pentyl, isopentyl, neopentyl, hexyl,
30 2-hexyl, 3-hexyl, 3-methylpentyl, vinyl, 2-pentene, cyclopropylmethyl, cyclopropyl, cyclohexylmethyl, cyclohexyl and propargyl. When reference is made herein to C1-C6 hydrocarbyl containing one or two double or triple bonds it is understood that at least two

carbons are present in the alkyl for one double or triple bond, and at least four carbons for two double or triple bonds.

By "nucleotide" is meant a heterocyclic nitrogenous base in N-glycosidic linkage with a phosphorylated sugar. Nucleotides are recognized in the art to include natural bases (standard), and modified bases well known in the art. Such bases are generally located at the 1' position of a nucleotide sugar moiety. Nucleotides generally comprise a base, sugar and a phosphate group. The nucleotides can be unmodified or modified at the sugar, phosphate and/or base moiety, (also referred to interchangeably as nucleotide analogs, modified nucleotides, non-natural nucleotides, non-standard nucleotides and other; see for example, Usman and McSwiggen, *supra*; Eckstein *et al.*, International PCT Publication No. WO 92/07065; Usman *et al.*, International PCT Publication No. WO 93/15187; Uhlman & Peyman, *supra* all are hereby incorporated by reference herein. There are several examples of modified nucleic acid bases known in the art as summarized by Limbach *et al.*, 1994, Nucleic Acids Res. 22, 2183. Some of the non-limiting examples of chemically modified and other natural nucleic acid bases that can be introduced into nucleic acids include, for example, inosine, purine, pyridin-4-one, pyridin-2-one, phenyl, pseudouracil, 2, 4, 6-trimethoxy benzene, 3-methyl uracil, dihydrouridine, naphthyl, aminophenyl, 5-alkylcytidines (*e.g.*, 5-methylcytidine), 5-alkyluridines (*e.g.*, ribothymidine), 5-halouridine (*e.g.*, 5-bromouridine) or 6-azapyrimidines or 6-alkylpyrimidines (*e.g.* 6-methyluridine), propyne, quesosine, 2-thiouridine, 4-thiouridine, wybutosine, wybutoxosine, 4-acetylcytidine, 5-(carboxyhydroxymethyl)uridine, 5'-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluridine, beta-D-galactosylqueosine, 1-methyladenosine, 1-methylinosine, 2,2-dimethylguanosine, 3-methylcytidine, 2-methyladenosine, 2-methylguanosine, N6-methyladenosine, 7-methylguanosine, 5-methoxyaminomethyl-2-thiouridine, 5-methylaminomethyluridine, 5-methylcarbonylmethyluridine, 5-methoxyuridine, 5-methyl-2-thiouridine, 2-methylthio-N6-isopentenyladenosine, beta-D-mannosylqueosine, uridine-5-oxyacetic acid, 2-thiocytidine, threonine derivatives and others (Burgin *et al.*, 1996, Biochemistry, 35, 14090; Uhlman & Peyman, *supra*). By "modified bases" in this aspect is meant nucleotide bases other than adenine, guanine, cytosine and uracil at 1' position or their equivalents; such bases can be used at any position, for example, within the catalytic core of an enzymatic nucleic acid molecule and/or in the substrate-binding regions of the nucleic acid molecule.

By "nucleoside" is meant a heterocyclic nitrogenous base in N-glycosidic linkage with a sugar. Nucleosides are recognized in the art to include natural bases (standard), and modified bases well known in the art. Such bases are generally located at the 1' position of a

nucleoside sugar moiety. Nucleosides generally comprise a base and sugar group. The nucleosides can be unmodified or modified at the sugar, and/or base moiety (also referred to interchangeably as nucleoside analogs, modified nucleosides, non-natural nucleosides, non-standard nucleosides and other; see for example, Usman and McSwiggen, *supra*; Eckstein *et al.*, International PCT Publication No. WO 92/07065; Usman *et al.*, International PCT Publication No. WO 93/15187; Uhlman & Peyman, *supra* all are hereby incorporated by reference herein). There are several examples of modified nucleic acid bases known in the art as summarized by Limbach *et al.*, 1994, *Nucleic Acids Res.* 22, 2183. Some of the non-limiting examples of chemically modified and other natural nucleic acid bases that can be introduced into nucleic acids include, inosine, purine, pyridin-4-one, pyridin-2-one, phenyl, pseudouracil, 2, 4, 6-trimethoxy benzene, 3-methyl uracil, dihydrouridine, naphthyl, aminophenyl, 5-alkylcytidines (e.g., 5-methylcytidine), 5-alkyluridines (e.g., ribothymidine), 5-halouridine (e.g., 5-bromouridine) or 6-azapyrimidines or 6-alkylpyrimidines (e.g. 6-methyluridine), propyne, quinosine, 2-thiouridine, 4-thiouridine, wybutosine, wybutoxosine, 4-acetylcytidine, 5-(carboxyhydroxymethyl)uridine, 5'-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluridine, beta-D-galactosylqueosine, 1-methyladenosine, 1-methylinosine, 2,2-dimethylguanosine, 3-methylcytidine, 2-methyladenosine, 2-methylguanosine, N6-methyladenosine, 7-methylguanosine, 5-methoxyaminomethyl-2-thiouridine, 5-methylaminomethyluridine, 5-methylcarbonylmethyluridine, 5-methoxyuridine, 5-methyl-2-thiouridine, 2-methylthio-N6-isopentenyladenosine, beta-D-mannosylqueosine, uridine-5-oxyacetic acid, 2-thiocytidine, threonine derivatives and others (Burgin *et al.*, 1996, *Biochemistry*, 35, 14090; Uhlman & Peyman, *supra*). By "modified bases" in this aspect is meant nucleoside bases other than adenine, guanine, cytosine and uracil at 1' position or their equivalents; such bases can be used at any position, for example, within the catalytic core of an enzymatic nucleic acid molecule and/or in the substrate-binding regions of the nucleic acid molecule.

In one embodiment, the invention features modified enzymatic nucleic acid molecules with phosphate backbone modifications comprising one or more phosphorothioate, phosphorodithioate, methylphosphonate, morpholino, amidate carbamate, carboxymethyl, acetamidate, polyamide, sulfonate, sulfonamide, sulfamate, formacetal, thioformacetal, and/or alkylsilyl, substitutions. For a review of oligonucleotide backbone modifications see Hunziker and Leumann, 1995, *Nucleic Acid Analogues: Synthesis and Properties*, in *Modern Synthetic Methods*, VCH, 331-417, and Mesmaeker *et al.*, 1994, *Novel Backbone Replacements for Oligonucleotides*, in *Carbohydrate Modifications in Antisense Research*, ACS, 24-39. These references are hereby incorporated by reference herein.

By "abasic" is meant sugar moieties lacking a base or having other chemical groups in place of a base at the 1' position, for example a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative (for more details see Wincott *et al.*, International PCT publication No. WO 97/26270).

- 5 By "unmodified nucleoside" is meant one of the bases adenine, cytosine, guanine, thymine, uracil joined to the 1' carbon of β -D-ribo-furanose.

By "modified nucleoside" is meant any nucleotide base which contains a modification in the chemical structure of an unmodified nucleotide base, sugar and/or phosphate.

- 10 In connection with 2'-modified nucleotides as described for the present invention, by "amino" is meant 2'-NH₂ or 2'-O-NH₂, which can be modified or unmodified. Such modified groups are described, for example, in Eckstein *et al.*, U.S. Patent 5,672,695 and Matulic-Adamic *et al.*, WO 98/28317, respectively, which are both incorporated by reference in their entireties.

- 15 Various modifications to nucleic acid (*e.g.*, DNAzyme) structure can be made to enhance the utility of these molecules. For example, such modifications can enhance shelf-life, half-life *in vitro*, stability, and ease of introduction of such oligonucleotides to the target site, including *e.g.*, enhancing penetration of cellular membranes and conferring the ability to recognize and bind to targeted cells.

- 20 Use of these molecules can lead to better treatment of the disease progression by affording the possibility of combination therapies (*e.g.*, multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of enzymatic nucleic acid molecules (including different enzymatic nucleic acid molecule motifs) and/or other chemical or biological molecules). The treatment of subjects with nucleic acid molecules can
25 also include combinations of different types of nucleic acid molecules. Therapies can be devised which include a mixture of enzymatic nucleic acid molecules (including different enzymatic nucleic acid molecule motifs), antisense and/or 2-5A chimera molecules to one or more targets to alleviate symptoms of a disease.

Administration of Nucleic Acid Molecules

- 30 Methods for the delivery of nucleic acid molecules are described in Akhtar *et al.*, 1992, *Trends Cell Bio.*, 2, 139; and *Delivery Strategies for Antisense Oligonucleotide Therapeutics*, ed. Akhtar, 1995, which are both incorporated herein by reference. Sullivan *et al.*, PCT WO

94/02595, further describes the general methods for delivery of enzymatic RNA molecules. These protocols can be utilized for the delivery of virtually any nucleic acid molecule. Nucleic acid molecules can be administered to cells by a variety of methods known to those familiar to the art, including, but not restricted to, encapsulation in liposomes, by iontophoresis, or by incorporation into other vehicles, such as hydrogels, cyclodextrins, biodegradable nanocapsules, and bioadhesive microspheres. Alternatively, the nucleic acid/vehicle combination is locally delivered by direct injection or by use of an infusion pump. Other routes of delivery include, but are not limited to oral (tablet or pill form) and/or intrathecal delivery (Gold, 1997, *Neuroscience*, 76, 1153-1158). Other approaches include the use of various transport and carrier systems, for example though the use of conjugates and biodegradable polymers. For a comprehensive review on drug delivery strategies including CNS delivery, see Ho *et al.*, 1999, *Curr. Opin. Mol. Ther.*, 1, 336-343 and Jain, *Drug Delivery Systems: Technologies and Commercial Opportunities*, Decision Resources, 1998 and Groothuis *et al.*, 1997, *J. NeuroVirol.*, 3, 387-400. More detailed descriptions of nucleic acid delivery and administration are provided in Sullivan *et al.*, *supra*, Draper *et al.*, PCT WO93/23569, Beigelman *et al.*, PCT WO99/05094, and Klimuk *et al.*, PCT WO99/04819, all of which have been incorporated by reference herein.

The molecules of the instant invention can be used as pharmaceutical agents. Pharmaceutical agents prevent, inhibit the occurrence, or treat (alleviate a symptom to some extent, preferably all of the symptoms) of a disease state in a subject.

The negatively charged polynucleotides of the invention can be administered (*e.g.*, RNA, DNA or protein) and introduced into a subject by any standard means described herein and known in the art, with or without stabilizers, buffers, and the like, to form a pharmaceutical composition. When it is desired to use a liposome delivery mechanism, standard protocols for formation of liposomes can be followed. The compositions of the present invention can also be formulated and used as tablets, capsules or elixirs for oral administration; suppositories for rectal administration; sterile solutions; suspensions for injectable administration; and the other compositions known in the art.

The present invention also includes pharmaceutically acceptable formulations of the compounds described. These formulations include salts of the above compounds, *e.g.*, acid addition salts, for example, salts of hydrochloric, hydrobromic, acetic acid, and benzene sulfonic acid.

A pharmacological composition or formulation refers to a composition or formulation in a form suitable for administration, *e.g.*, systemic administration, into a cell or subject,

preferably a human. Suitable forms, in part, depend upon the use or the route of entry, for example oral, transdermal, or by injection. Such forms should not prevent the composition or formulation from reaching a target cell (*i.e.*, a cell to which the negatively charged polymer is desired to be delivered to). For example, pharmacological compositions injected into the blood stream should be soluble. Other factors are known in the art, and include considerations such as toxicity and forms which prevent the composition or formulation from exerting its effect.

By "systemic administration" is meant *in vivo* systemic absorption or accumulation of drugs in the blood stream followed by distribution throughout the entire body. Administration routes which lead to systemic absorption include, without limitations: intravenous, subcutaneous, intraperitoneal, inhalation, oral, intrapulmonary and intramuscular. Each of these administration routes expose the desired negatively charged polymers, *e.g.*, nucleic acids, to an accessible diseased tissue. The rate of entry of a drug into the circulation has been shown to be a function of molecular weight or size. The use of a liposome or other drug carrier comprising the compounds of the instant invention can potentially localize the drug, for example, in certain tissue types, such as the tissues of the reticular endothelial system (RES). A liposome formulation that can facilitate the association of drug with the surface of cells, such as, lymphocytes and macrophages is also useful. This approach can provide enhanced delivery of the drug to target cells by taking advantage of the specificity of macrophage and lymphocyte immune recognition of abnormal cells, such as cancer cells.

By pharmaceutically acceptable formulation is meant, a composition or formulation that allows for the effective distribution of the nucleic acid molecules of the instant invention in the physical location most suitable for their desired activity. Non-limiting examples of agents suitable for formulation with the nucleic acid molecules of the instant invention include: PEG conjugated nucleic acids, phospholipid conjugated nucleic acids, nucleic acids containing lipophilic moieties, phosphorothioates, P-glycoprotein inhibitors (such as Pluronic P85) which can enhance entry of drugs into various tissues, for example the CNS (Jolliet-Riant and Tillement, 1999, *Fundam. Clin. Pharmacol.*, 13, 16-26); biodegradable polymers, such as poly (DL-lactide-coglycolide) microspheres for sustained release delivery after implantation (Emerich, DF *et al*, 1999, *Cell Transplant*, 8, 47-58) Alkermes, Inc. Cambridge, MA; and loaded nanoparticles, such as those made of polybutylcyanoacrylate, which can deliver drugs across the blood brain barrier and can alter neuronal uptake mechanisms (*Prog Neuropsychopharmacol Biol Psychiatry*, 23, 941-949, 1999). Other non-limiting examples of delivery strategies, including CNS delivery of the nucleic acid molecules of the instant

invention include material described in Boado *et al.*, 1998, *J. Pharm. Sci.*, 87, 1308-1315; Tyler *et al.*, 1999, *FEBS Lett.*, 421, 280-284; Pardridge *et al.*, 1995, *PNAS USA.*, 92, 5592-5596; Boado, 1995, *Adv. Drug Delivery Rev.*, 15, 73-107; Aldrian-Herrada *et al.*, 1998, *Nucleic Acids Res.*, 26, 4910-4916; and Tyler *et al.*, 1999, *PNAS USA.*, 96, 7053-7058. All these references are hereby incorporated herein by reference.

The invention also features the use of the composition comprising surface-modified liposomes containing poly (ethylene glycol) lipids (PEG-modified, or long-circulating liposomes or stealth liposomes). Nucleic acid molecules of the invention can also comprise covalently attached PEG molecules of various molecular weights. These formulations offer a method for increasing the accumulation of drugs in target tissues. This class of drug carriers resists opsonization and elimination by the mononuclear phagocytic system (MPS or RES), thereby enabling longer blood circulation times and enhanced tissue exposure for the encapsulated drug (Lasic *et al. Chem. Rev.* 1995, 95, 2601-2627; Ishiwata *et al., Chem. Pharm. Bull.* 1995, 43, 1005-1011). Such liposomes have been shown to accumulate selectively in tumors, presumably by extravasation and capture in the neovascularized target tissues (Lasic *et al., Science* 1995, 267, 1275-1276; Oku *et al., 1995, Biochim. Biophys. Acta*, 1238, 86-90). The long-circulating liposomes enhance the pharmacokinetics and pharmacodynamics of DNA and RNA, particularly compared to conventional cationic liposomes, which are known to accumulate in tissues of the MPS (Liu *et al., J. Biol. Chem.* 1995, 270, 24864-24870; Choi *et al.*, International PCT Publication No. WO 96/10391; Ansell *et al.*, International PCT Publication No. WO 96/10390; Holland *et al.*, International PCT Publication No. WO 96/10392; all of which are incorporated by reference herein). Long-circulating liposomes are also likely to protect drugs from nuclease degradation to a greater extent compared to cationic liposomes, based on their ability to avoid accumulation in metabolically aggressive MPS tissues such as the liver and spleen. All of these references are incorporated by reference herein.

The present invention also includes compositions prepared for storage or administration that include a pharmaceutically effective amount of the desired compounds in a pharmaceutically acceptable carrier or diluent. Acceptable carriers or diluents for therapeutic use are well known in the pharmaceutical art, and are described, for example, in *Remington's Pharmaceutical Sciences*, Mack Publishing Co. (A.R. Gennaro edit. 1985), hereby incorporated by reference herein. For example, preservatives, stabilizers, dyes and flavoring agents can be provided. These include sodium benzoate, sorbic acid and esters of *p*-hydroxybenzoic acid. In addition, antioxidants and suspending agents can be used.

A pharmaceutically effective dose is that dose required to prevent, inhibit the occurrence, or treat (alleviate a symptom to some extent, preferably all of the symptoms) of a disease state. The pharmaceutically effective dose depends on the type of disease, the composition used, the route of administration, the type of mammal being treated, the physical characteristics of the specific mammal under consideration, concurrent medication, and other factors which those skilled in the medical arts will recognize. Generally, an amount between 0.1 mg/kg and 100 mg/kg body weight/day of active ingredients is administered dependent upon potency of the negatively charged polymer.

The nucleic acid molecules of the invention and formulations thereof can be administered orally, topically, parenterally, by inhalation or spray, or rectally in dosage unit formulations containing conventional non-toxic pharmaceutically acceptable carriers, adjuvants and/or vehicles. The term parenteral as used herein includes percutaneous, subcutaneous, intravascular (e.g., intravenous), intramuscular, or intrathecal injection or infusion techniques and the like. In addition, there is provided a pharmaceutical formulation comprising a nucleic acid molecule of the invention and a pharmaceutically acceptable carrier. One or more nucleic acid molecules of the invention can be present in association with one or more non-toxic pharmaceutically acceptable carriers and/or diluents and/or adjuvants, and if desired other active ingredients. The pharmaceutical compositions containing nucleic acid molecules of the invention can be in a form suitable for oral use, for example, as tablets, troches, lozenges, aqueous or oily suspensions, dispersible powders or granules, emulsion, hard or soft capsules, or syrups or elixirs.

Compositions intended for oral use can be prepared according to any method known to the art for the manufacture of pharmaceutical compositions and such compositions can contain one or more such sweetening agents, flavoring agents, coloring agents or preservative agents in order to provide pharmaceutically elegant and palatable preparations. Tablets contain the active ingredient in admixture with non-toxic pharmaceutically acceptable excipients that are suitable for the manufacture of tablets. These excipients can be, for example, inert diluents, such as calcium carbonate, sodium carbonate, lactose, calcium phosphate or sodium phosphate; granulating and disintegrating agents, for example, corn starch, or alginic acid; binding agents, for example starch, gelatin or acacia, and lubricating agents, for example magnesium stearate, stearic acid or talc. The tablets can be uncoated or they can be coated by known techniques. In some cases such coatings can be prepared by known techniques to delay disintegration and absorption in the gastrointestinal tract and thereby provide a sustained action over a longer period. For example, a time delay material such as glyceryl monostearate or glyceryl distearate can be employed.

Formulations for oral use can also be presented as hard gelatin capsules wherein the active ingredient is mixed with an inert solid diluent, for example, calcium carbonate, calcium phosphate or kaolin, or as soft gelatin capsules wherein the active ingredient is mixed with water or an oil medium, for example peanut oil, liquid paraffin or olive oil.

5 Aqueous suspensions contain the active materials in admixture with excipients suitable for the manufacture of aqueous suspensions. Such excipients are suspending agents, for example, sodium carboxymethylcellulose, methylcellulose, hydropropyl-methylcellulose, sodium alginate, polyvinylpyrrolidone, gum tragacanth and gum acacia; dispersing or wetting
10 agents can be a naturally-occurring phosphatide, for example, lecithin, or condensation products of an alkylene oxide with fatty acids, for example polyoxyethylene stearate, or condensation products of ethylene oxide with long chain aliphatic alcohols, for example heptadecaethyleneoxycetanol, or condensation products of ethylene oxide with partial esters derived from fatty acids and a hexitol such as polyoxyethylene sorbitol monooleate, or
15 condensation products of ethylene oxide with partial esters derived from fatty acids and hexitol anhydrides, for example polyethylene sorbitan monooleate. The aqueous suspensions can also contain one or more preservatives, for example, ethyl, or n-propyl p-hydroxybenzoate, one or more coloring agents, one or more flavoring agents, and one or more sweetening agents, such as sucrose or saccharin.

20 Oily suspensions can be formulated by suspending the active ingredients in a vegetable oil, for example arachis oil, olive oil, sesame oil or coconut oil, or in a mineral oil such as liquid paraffin. The oily suspensions can contain a thickening agent, for example beeswax, hard paraffin or cetyl alcohol. Sweetening agents and flavoring agents can be added to provide palatable oral preparations. These compositions can be preserved by the addition of an anti-oxidant such as ascorbic acid.

25 Dispersible powders and granules suitable for preparation of an aqueous suspension by the addition of water provide the active ingredient in admixture with a dispersing or wetting agent, suspending agent and one or more preservatives. Suitable dispersing or wetting agents or suspending agents are exemplified by those already mentioned above. Additional excipients, for example sweetening, flavoring and coloring agents, can also be present.

30 Pharmaceutical compositions of the invention can also be in the form of oil-in-water emulsions. The oily phase can be a vegetable oil or a mineral oil or mixtures of these. Suitable emulsifying agents can be naturally-occurring gums, for example gum acacia or gum tragacanth, naturally-occurring phosphatides, for example soy bean, lecithin, and esters or

partial esters derived from fatty acids and hexitol, anhydrides, for example, sorbitan monooleate, and condensation products of the said partial esters with ethylene oxide, for example polyoxyethylene sorbitan monooleate. The emulsions can also contain sweetening and flavoring agents.

5 Syrups and elixirs can be formulated with sweetening agents, for example glycerol, propylene glycol, sorbitol, glucose or sucrose. Such formulations can also contain a demulcent, a preservative and flavoring and coloring agents. The pharmaceutical compositions can be in the form of a sterile injectable aqueous or oleaginous suspension. This suspension can be formulated according to the known art using those suitable dispersing
10 or wetting agents and suspending agents that have been mentioned above. The sterile injectable preparation can also be a sterile injectable solution or suspension in a non-toxic parentally acceptable diluent or solvent, for example as a solution in 1,3-butanediol. Among the acceptable vehicles and solvents that can be employed are water, Ringer's solution and isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed
15 as a solvent or suspending medium. For this purpose any bland fixed oil can be employed including synthetic mono-or diglycerides. In addition, fatty acids such as oleic acid find use in the preparation of injectables.

 The nucleic acid molecules of the invention can also be administered in the form of suppositories, *e.g.*, for rectal administration of the drug. These compositions can be prepared
20 by mixing the drug with a suitable non-irritating excipient that is solid at ordinary temperatures but liquid at the rectal temperature and will therefore melt in the rectum to release the drug. Such materials include cocoa butter and polyethylene glycols.

 Nucleic acid molecules of the invention can be administered parenterally in a sterile medium. The drug, depending on the vehicle and concentration used, can either be suspended
25 or dissolved in the vehicle. Advantageously, adjuvants such as local anesthetics, preservatives and buffering agents can be dissolved in the vehicle.

 Dosage levels of the order of from about 0.1 mg to about 140 mg per kilogram of body weight per day are useful in the treatment of the above-indicated conditions (about 0.5 mg to about 7 g per patient or subject per day). The amount of active ingredient that can be
30 combined with the carrier materials to produce a single dosage form varies depending upon the host treated and the particular mode of administration. Dosage unit forms generally contain between from about 1 mg to about 500 mg of an active ingredient.

It is understood that the specific dose level for any particular patient or subject depends upon a variety of factors including the activity of the specific compound employed, the age, body weight, general health, sex, diet, time of administration, route of administration, and rate of excretion, drug combination and the severity of the particular disease undergoing therapy.

5 For administration to non-human animals, the composition can also be added to the animal feed or drinking water. It can be convenient to formulate the animal feed and drinking water compositions so that the animal takes in a therapeutically appropriate quantity of the composition along with its diet. It can also be convenient to present the composition as a premix for addition to the feed or drinking water.

10 The nucleic acid molecules of the present invention can also be administered to a patient or subject in combination with other therapeutic compounds to increase the overall therapeutic effect. The use of multiple compounds to treat an indication can increase the beneficial effects while reducing the presence of side effects.

In another aspect of the invention, nucleic acid molecules of the present invention are preferably expressed from transcription units (see for example Couture *et al.*, 1996, *TIG.*, 12, 510, Skillern *et al.*, International PCT Publication No. WO 00/22113, Conrad, International PCT Publication No. WO 00/22114, and Conrad, US 6,054,299) inserted into DNA or RNA vectors. The recombinant vectors are preferably DNA plasmids or viral vectors. Enzymatic nucleic acid expressing viral vectors can be constructed based on, but not limited to, adeno-associated virus, retrovirus, adenovirus, or alphavirus. Preferably, the recombinant vectors capable of expressing the nucleic acid molecules are delivered as described above, and persist in target cells. Alternatively, viral vectors can be used that provide for transient expression of nucleic acid molecules. Such vectors can be repeatedly administered as necessary. Once expressed, the nucleic acid molecule binds to the target mRNA. Delivery of nucleic acid molecule expressing vectors can be systemic, such as by intravenous or intra-muscular administration, by administration to target cells ex-planted from the subject followed by reintroduction into the subject, or by any other means that would allow for introduction into the desired target cell (for a review see Couture *et al.*, 1996, *TIG.*, 12, 510).

30 One aspect of the invention features an expression vector comprising a nucleic acid sequence encoding at least one of the nucleic acid molecules of the instant invention. The nucleic acid sequence encoding the nucleic acid molecule of the instant invention is operably linked in a manner that allows expression of that nucleic acid molecule.

Another aspect the invention features an expression vector comprising nucleic acid sequence encoding at least one of the nucleic acid molecules of the invention, in a manner which allows expression of that nucleic acid molecule. The expression vector comprises in one embodiment; a) a transcription initiation region; b) a transcription termination region; c) a nucleic acid sequence encoding at least one said nucleic acid molecule; and wherein said sequence is operably linked to said initiation region and said termination region, in a manner that allows expression and/or delivery of said nucleic acid molecule.

In another embodiment, the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an open reading frame; d) a nucleic acid sequence encoding at least one said nucleic acid molecule, wherein said sequence is operably linked to the 3'-end of said open reading frame; and wherein said sequence is operably linked to said initiation region, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule. In yet another embodiment the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an intron; d) a nucleic acid sequence encoding at least one said nucleic acid molecule; and wherein said sequence is operably linked to said initiation region, said intron and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule.

In another embodiment, the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an intron; d) an open reading frame; e) a nucleic acid sequence encoding at least one said nucleic acid molecule, wherein said sequence is operably linked to the 3'-end of said open reading frame; and wherein said sequence is operably linked to said initiation region, said intron, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule.

Examples

The following are non-limiting examples showing the selection, isolation, synthesis and activity of nucleic acids of the instant invention.

Example 1: Identification of Potential Target Sites in Human Ras RNA

The sequence of human Ras genes were screened for accessible sites using a computer-folding algorithm. Regions of the RNA that do not form secondary folding structures and contain potential enzymatic nucleic acid molecule and/or antisense binding/cleavage sites

were identified. The sequences of K-Ras and H-Ras binding/cleavage sites are shown in **Tables II and III**.

Example 2: Selection of Enzymatic Nucleic Acid Cleavage Sites in Human Ras RNA

Enzymatic nucleic acid molecule target sites were chosen by analyzing sequences of Human K-Ras and H-Ras (for example, Genbank accession Nos: NM_004985 and NM_005343 respectively) and prioritizing the sites on the basis of folding. Enzymatic nucleic acid molecules were designed that can bind each target and were individually analyzed by computer folding (Christoffersen *et al.*, 1994 *J. Mol. Struct. Theochem*, 311, 273; Jacger *et al.*, 1989, *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether the enzymatic nucleic acid molecule sequences fold into the appropriate secondary structure. Those enzymatic nucleic acid molecules with unfavorable intramolecular interactions between the binding arms and the catalytic core are eliminated from consideration. As noted below, varying binding arm lengths can be chosen to optimize activity. Generally, at least 5 bases on each arm are able to bind to, or otherwise interact with, the target RNA.

Example 3: Chemical Synthesis and Purification of Enzymatic Nucleic Acid Molecules for Efficient Cleavage and/or blocking of Ras RNA

DNAzyme molecules are designed to anneal to various sites in the RNA message. The binding arms of the DNAzyme molecules are complementary to the target site sequences described above. The DNAzymes were chemically synthesized. The method of synthesis used followed the procedure for nucleic acid synthesis as described herein and in Usman *et al.*, (1987 *J. Am. Chem. Soc.*, 109, 7845), Scaringe *et al.*, (1990 *Nucleic Acids Res.*, 18, 5433) and Wincott *et al.*, *supra*, and made use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. The average stepwise coupling yields were typically >98%. The sequences of the chemically synthesized DNAzyme molecules used in this study are shown below in **Tables II and III**.

Example 4: DNAzyme Cleavage of Ras RNA Target *in vitro*

DNAzymes targeted to the human K-Ras and H-Ras RNA are designed and synthesized as described above. These enzymatic nucleic acid molecules can be tested for cleavage activity *in vitro*, for example, using the following procedure. The target sequences and the nucleotide location within the K-Ras and H-Ras RNA are given in **Tables II and III** respectively.

Cleavage Reactions:

DNAzymes and substrates were synthesized in 96-well format using 0.2 μ mol scale. Substrates were 5'-³²P labeled and gel purified using 7.5% polyacrylamide gels, and eluting into water. Assays were done by combining trace substrate with 500nM DNAzyme or greater, and initiated by adding final concentrations of 40mM Mg⁺², and 50mM Tris-Cl pH 8.0. For each DNAzyme/substrate combination a control reaction was done to ensure cleavage was not the result of non-specific substrate degradation. A single three hour time point was taken and run on a 15% polyacrylamide gel to assess cleavage activity. Gels were dried and scanned using a Molecular Dynamics Phosphorimager and quantified using Molecular Dynamics ImageQuant software. Percent cleaved was determined by dividing values for cleaved substrate bands by full-length (uncleaved) values plus cleaved values and multiplying by 100 (%cleaved=[C/(U+C)]*100).

Example 5: DNAzyme Cleavage of Ras RNA Target *in vivo*

Cell Culture

Wickstrom, 2001, *Mol. Biotechnol.*, 18, 35-35, describes a cell culture system in which antisense oligonucleotides targeting H-Ras were studied in transformed mouse cells that form solid tumors. Treatment of cells with antisense targeting H-Ras resulted in the sequence specific and dose dependent inhibition of H-Ras expression. In this study, it was determined that antisense targeting the first intron region of H-Ras were more effective than antisense targeting the initiation codon region.

Kita *et al.*, 1999, *Int. J. Cancer*, 80, 553-558, describes the growth inhibition of human pancreatic cancer cell lines by antisense oligonucleotides specific to mutated K-Ras genes. Antisense oligonucleotides were transfected to the transformed cells using liposomes. Cellular proliferation, K-Ras mRNA expression, and K-Ras protein synthesis were all evaluated as endpoints. Sato *et al.*, 2000, *Cancer Lett.*, 155, 153-161, describes another human pancreatic cancer cell line, HOR-P1, that is characterized by high angiogenic activity and metastatic potential. Genetic and molecular analysis of this cell line revealed both increased telomerase activity and a mutation in the K-Ras oncogene.

A variety of endpoints have been used in cell culture models to look at Ras-mediated effects after treatment with anti-Ras agents. Phenotypic endpoints include inhibition of cell proliferation, RNA expression, and reduction of Ras protein expression. Because Ras oncogene mutations are directly associated with increased proliferation of certain tumor cells,

a proliferation endpoint for cell culture assays is preferably used as the primary screen. There are several methods by which this endpoint can be measured. Following treatment of cells with DNAzymes, cells are allowed to grow (typically 5 days) after which either the cell viability, the incorporation of [³H] thymidine into cellular DNA and/or the cell density can be measured. The assay of cell density is done in a 96-well format using commercially available fluorescent nucleic acid stains (such as Syto® 13 or CyQuant®). As a secondary, confirmatory endpoint a DNAzyme-mediated decrease in the level of Ras protein expression is evaluated using a Ras-specific ELISA.

Animal Models

Evaluating the efficacy of anti-Ras agents in animal models is an important prerequisite to human clinical trials. As in cell culture models, the most Ras sensitive mouse tumor xenografts are those derived from cancer cells that express mutant Ras proteins. Nude mice bearing H-Ras transformed bladder cancer cell xenografts were sensitive to an anti-Ras antisense nucleic acid, resulting in an 80% inhibition of tumor growth after a 31 day treatment period (Wickstrom, 2001, *Mol. Biotechnol.*, 18, 35-35). Zhang *et al.*, 2000, *Gene Ther.*, 7, 2041, describes an anti-K-Ras ribozyme adenoviral vector (KRbz-ADV) targeting a K-Ras mutant (K-Ras codon 12 GGT to GTT; H441 and H1725 cells respectively). Non-small cell lung cancer cells (NSCLC H441 and H1725 cells) that express the mutant K-Ras protein were used in nude mouse xenografts compared to NSCLC H1650 cells that lack the relevant mutation. Pre-treatment with KRbz-ADV completely abrogated engraftment of both H441 and H1725 cells and compared to 100% engraftment and tumor growth in animals that received untreated tumor cells or a control vector. The above studies provide proof that inhibition of Ras expression by anti-Ras agents causes inhibition of tumor growth in animals. Anti-Ras DNAzymes chosen from *in vitro* assays are further tested in similar mouse xenograft models. Active DNAzymes are subsequently tested in combination with standard chemotherapies.

Indications

Particular degenerative and disease states that are associated with Ras expression modulation include but are not limited to cancer, for example lung cancer, colorectal cancer, bladder cancer, pancreatic cancer, breast cancer, prostate cancer and/or any other diseases or conditions that are related to or will respond to the levels of Ras in a cell or tissue, alone or in combination with other therapies.

The present body of knowledge in Ras research indicates the need for methods to assay Ras activity and for compounds that can regulate Ras expression for research, diagnostic, and therapeutic use.

The use of monoclonal antibodies, chemotherapy, radiation therapy, and analgesics, are all non-limiting examples of methods that can be combined with or used in conjunction with the nucleic acid molecules (e.g. DNazymes) of the instant invention. Common chemotherapies that can be combined with nucleic acid molecules of the instant invention include various combinations of cytotoxic drugs to kill cancer cells. These drugs include but are not limited to paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, vinorelbine etc. Those skilled in the art will recognize that other drug compounds and therapies can be similarly be readily combined with the nucleic acid molecules of the instant invention (e.g. DNzyme molecules) are hence within the scope of the instant invention.

Diagnostic uses

The nucleic acid molecules of this invention (e.g., enzymatic nucleic acid molecules) are used as diagnostic tools to examine genetic drift and mutations within diseased cells or to detect the presence of Ras RNA in a cell. The close relationship between enzymatic nucleic acid molecule activity and the structure of the target RNA allows the detection of mutations in any region of the molecule that alters the base-pairing and three-dimensional structure of the target RNA. Using multiple enzymatic nucleic acid molecules described in this invention, one maps nucleotide changes which are important to RNA structure and function *in vitro*, as well as in cells and tissues. Cleavage of target RNAs with enzymatic nucleic acid molecules are used to inhibit gene expression and define the role (essentially) of specified gene products in the progression of disease. In this manner, other genetic targets are defined as important mediators of the disease. These experiments lead to better treatment of the disease progression by affording the possibility of combinational therapies (e.g., multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of enzymatic nucleic acid molecules and/or other chemical or biological molecules). Other *in vitro* uses of enzymatic nucleic acid molecules of this invention are known in the art, and include detection of the presence of mRNAs associated with Ras-related condition. Such RNA is detected by determining the presence of a cleavage product after treatment with an enzymatic nucleic acid molecule using standard methodology.

In a specific example, enzymatic nucleic acid molecules that cleave only wild-type or mutant forms of the target RNA are used for the assay. The first enzymatic nucleic acid molecule is used to identify wild-type RNA present in the sample and the second enzymatic nucleic acid molecule is used to identify mutant RNA in the sample. As reaction controls, synthetic substrates of both wild-type and mutant RNA are cleaved by both enzymatic nucleic acid molecules to demonstrate the relative enzymatic nucleic acid molecule efficiencies in the reactions and the absence of cleavage of the "non-targeted" RNA species. The cleavage products from the synthetic substrates also serve to generate size markers for the analysis of wild-type and mutant RNAs in the sample population. Thus each analysis requires two enzymatic nucleic acid molecules, two substrates and one unknown sample which is combined into six reactions. The presence of cleavage products is determined using an RNase protection assay so that full-length and cleavage fragments of each RNA can be analyzed in one lane of a polyacrylamide gel. It is not absolutely required to quantify the results to gain insight into the expression of mutant RNAs and putative risk of the desired phenotypic changes in target cells. The expression of mRNA whose protein product is implicated in the development of the phenotype (*i.e.*, Ras) is adequate to establish risk. If probes of comparable specific activity are used for both transcripts, then a qualitative comparison of RNA levels will be adequate and will decrease the cost of the initial diagnosis. Higher mutant form to wild-type ratios are correlated with higher risk whether RNA levels are compared qualitatively or quantitatively. The use of enzymatic nucleic acid molecules in diagnostic applications contemplated by the instant invention is described, for example, in George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, International PCT publication No. WO 99/29842.

Example 6: Identification of Potential Target Sites in Human HIV RNA

The sequence of human HIV genes are screened for accessible sites using a computer-folding algorithm. Regions of the RNA that do not form secondary folding structures and contained potential enzymatic nucleic acid molecule and/or antisense binding/cleavage sites are identified. The sequences of these binding/cleavage sites are shown in Tables VI to XI.

Example 6: Selection of Enzymatic Nucleic Acid Cleavage Sites in Human HIV RNA

Enzymatic nucleic acid molecule target sites were chosen by analyzing sequences of Human HIV (Genbank accession No: NM_005228) and prioritizing the sites on the basis of

folding. Enzymatic nucleic acid molecules were designed that can bind each target and are individually analyzed by computer folding (Christoffersen *et al.*, 1994 *J. Mol. Struct. Theochem*, 311, 273; Jaeger *et al.*, 1989, *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether the enzymatic nucleic acid molecule sequences fold into the appropriate secondary structure. Those enzymatic nucleic acid molecules with unfavorable intramolecular interactions between the binding arms and the catalytic core were eliminated from consideration. As noted below, varying binding arm lengths can be chosen to optimize activity. Generally, at least 5 bases on each arm are able to bind to, or otherwise interact with, the target RNA.

10 Example 8: Chemical Synthesis and Purification of Ribozymes and Antisense for Efficient Cleavage and/or blocking of HIV Activity

Enzymatic nucleic acid molecules and antisense constructs are designed to anneal to various sites in the RNA message. The binding arms of the enzymatic nucleic acid molecules are complementary to the target site sequences described above, while the antisense constructs are fully complementary to the target site sequences described above. The enzymatic nucleic acid molecules and antisense constructs were chemically synthesized. The method of synthesis used followed the procedure for normal RNA synthesis as described above and in Usman *et al.*, (1987 *J. Am. Chem. Soc.*, 109, 7845), Scaringe *et al.*, (1990 *Nucleic Acids Res.*, 18, 5433) and Wincott *et al.*, *supra*, and made use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. The average stepwise coupling yields were typically >98%.

Enzymatic nucleic acid molecules and antisense constructs are also synthesized from DNA templates using bacteriophage T7 RNA polymerase (Milligan and Uhlenbeck, 1989, *Methods Enzymol.* 180, 51). Enzymatic nucleic acid molecules and antisense constructs are purified by gel electrophoresis using general methods or are purified by high pressure liquid chromatography (HPLC; See Wincott *et al.*, *supra*; the totality of which is hereby incorporated herein by reference) and are resuspended in water. The sequences of the chemically synthesized enzymatic nucleic acid molecules used in this study are shown below in Table XI. The sequences of the chemically synthesized antisense constructs used in this study are complementary sequences to the Substrate sequences shown below as in Tables VI to XI.

Example 8: Enzymatic nucleic acid molecule Cleavage of HIV RNA Target *in vitro*

Enzymatic nucleic acid molecules targeted to the human HIV RNA are designed and synthesized as described above. These enzymatic nucleic acid molecules are tested for cleavage activity *in vitro*, for example, using the following procedure. The target sequences and the nucleotide location within the HIV RNA are given in Tables VI to XI.

5 *Cleavage Reactions:* Full-length or partially full-length, internally-labeled target RNA for enzymatic nucleic acid molecule cleavage assay is prepared by *in vitro* transcription in the presence of [α - 32 P] CTP, passed over a G 50 Sephadex column by spin chromatography and used as substrate RNA without further purification. Alternately, substrates are 5'- 32 P-end labeled using T4 polynucleotide kinase enzyme. Assays are performed by pre-warming a 2X
10 concentration of purified enzymatic nucleic acid molecule in enzymatic nucleic acid molecule cleavage buffer (50 mM Tris-HCl, pH 7.5 at 37°C, 10 mM MgCl₂) and the cleavage reaction was initiated by adding the 2X enzymatic nucleic acid molecule mix to an equal volume of substrate RNA (maximum of 1-5 nM) that was also pre-warmed in cleavage buffer. As an
15 initial screen, assays are carried out for 1 hour at 37°C using a final concentration of either 40 nM or 1 mM enzymatic nucleic acid molecule, *i.e.*, enzymatic nucleic acid molecule excess. The reaction is quenched by the addition of an equal volume of 95% formamide, 20 mM EDTA, 0.05% bromophenol blue and 0.05% xylene cyanol after which the sample is heated to 95°C for 2 minutes, quick chilled and loaded onto a denaturing polyacrylamide gel. Substrate RNA and the specific RNA cleavage products generated by enzymatic nucleic acid
20 molecule cleavage are visualized on an autoradiograph of the gel. The percentage of cleavage is determined by Phosphor Imager[®] quantitation of bands representing the intact substrate and the cleavage products.

Indications

25 Particular degenerative and disease states that can be associated with HIV expression modulation include but are not limited to acquired immunodeficiency disease (AIDS) and related diseases and conditions, including but not limited to Kaposi's sarcoma, lymphoma, cervical cancer, squamous cell carcinoma, cardiac myopathy, rheumatic diseases, and opportunistic infection, for example Pneumocystis carinii, Cytomegalovirus, Herpes simplex, Mycobacteria, Cryptococcus, Toxoplasma, Progressive multifocal leucoencephalopathy
30 (Papovavirus), Mycobacteria, Aspergillus, Cryptococcus, Candida, Cryptosporidium, Isospora belli, Microsporidia and any other diseases or conditions that are related to or will respond to the levels of HIV in a cell or tissue, alone or in combination with other therapies

The present body of knowledge in HIV research indicates the need for methods to assay HIV activity and for compounds that can regulate HIV expression for research, diagnostic, and therapeutic use.

The use of antiviral compounds, monoclonal antibodies, chemotherapy, radiation therapy, analgesics, and/or anti-inflammatory compounds, are all non-limiting examples of a methods that can be combined with or used in conjunction with the nucleic acid molecules (e.g. ribozymes and antisense molecules) of the instant invention. Examples of antiviral compounds that can be used in conjunction with the nucleic acid molecules of the invention include but are not limited to AZT (also known as zidovudine or ZDV), ddC (zalcitabine), ddI (dideoxyinosine), d4T (stavudine), and 3TC (lamivudine) Ribavirin, delvaridine (Rescriptor), nevirapine (Viramune), efavirenz (Sustiva), ritonavir (Norvir), saquinivir (Invirase), indinavir (Crixivan), amprenivir (Agenerase), nelfinavir (Viracept), and/or lopinavir (Kaletra). Common chemotherapies that can be combined with nucleic acid molecules of the instant invention include various combinations of cytotoxic drugs to kill cancer cells. These drugs include but are not limited to paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, vinorelbine etc. Those skilled in the art will recognize that other drug compounds and therapies can be similarly be readily combined with the nucleic acid molecules of the instant invention (e.g. ribozymes and antisense molecules) are hence within the scope of the instant invention.

Diagnostic uses

The nucleic acid molecules of this invention (e.g., enzymatic nucleic acid molecules) are used as diagnostic tools to examine genetic drift and mutations within diseased cells or to detect the presence of HIV RNA in a cell. The close relationship between enzymatic nucleic acid molecule activity and the structure of the target RNA allows the detection of mutations in any region of the molecule which alters the base-pairing and three-dimensional structure of the target RNA. Using multiple enzymatic nucleic acid molecules described in this invention, one maps nucleotide changes which are important to RNA structure and function *in vitro*, as well as in cells and tissues. Cleavage of target RNAs with enzymatic nucleic acid molecules are used to inhibit gene expression and define the role (essentially) of specified gene products in the progression of disease. In this manner, other genetic targets are defined as important mediators of the disease. These experiments lead to better treatment of the disease progression by affording the possibility of combinational therapies (e.g., multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of

enzymatic nucleic acid molecules and/or other chemical or biological molecules). Other *in vitro* uses of enzymatic nucleic acid molecules of this invention are well known in the art, and include detection of the presence of mRNAs associated with HIV-related condition. Such RNA is detected by determining the presence of a cleavage product after treatment with an enzymatic nucleic acid molecule using standard methodology.

In a specific example, enzymatic nucleic acid molecules which cleave only wild-type or mutant forms of the target RNA are used for the assay. The first enzymatic nucleic acid molecule is used to identify wild-type RNA present in the sample and the second enzymatic nucleic acid molecule is used to identify mutant RNA in the sample. As reaction controls, synthetic substrates of both wild-type and mutant RNA are cleaved by both enzymatic nucleic acid molecules to demonstrate the relative enzymatic nucleic acid molecule efficiencies in the reactions and the absence of cleavage of the "non-targeted" RNA species. The cleavage products from the synthetic substrates also serve to generate size markers for the analysis of wild-type and mutant RNAs in the sample population. Thus each analysis requires two enzymatic nucleic acid molecules, two substrates and one unknown sample which is combined into six reactions. The presence of cleavage products is determined using an RNase protection assay so that full-length and cleavage fragments of each RNA can be analyzed in one lane of a polyacrylamide gel. It is not absolutely required to quantify the results to gain insight into the expression of mutant RNAs and putative risk of the desired phenotypic changes in target cells. The expression of mRNA whose protein product is implicated in the development of the phenotype (*i.e.*, HIV) is adequate to establish risk. If probes of comparable specific activity are used for both transcripts, then a qualitative comparison of RNA levels will be adequate and will decrease the cost of the initial diagnosis. Higher mutant form to wild-type ratios are correlated with higher risk whether RNA levels are compared qualitatively or quantitatively. The use of enzymatic nucleic acid molecules in diagnostic applications contemplated by the instant invention is more fully described in George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, International PCT publication No. WO 99/29842.

Example 10: Identification of Potential Target Sites in Human HER2 RNA

The sequence of human HER2 genes were screened for accessible sites using a computer-folding algorithm. Regions of the RNA that do not form secondary folding structures and contained potential enzymatic nucleic acid molecule and/or antisense

binding/cleavage sites were identified. The sequences of these binding/cleavage sites are shown in Tables IV and V.

Example 10: Selection of Enzymatic Nucleic Acid Cleavage Sites in Human HER2 RNA

Enzymatic nucleic acid molecule target sites were chosen by analyzing sequences of Human HER2 (Genbank accession No: X03363) and prioritizing the sites on the basis of folding. Enzymatic nucleic acid molecules were designed that can bind each target and are individually analyzed by computer folding (Christoffersen *et al.*, 1994 *J. Mol. Struct. Theochem*, 311, 273; Jaeger *et al.*, 1989, *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether the enzymatic nucleic acid molecule sequences fold into the appropriate secondary structure. Those enzymatic nucleic acid molecules with unfavorable intramolecular interactions between the binding arms and the catalytic core were eliminated from consideration. As noted below, variable binding arm lengths are chosen to optimize activity. Generally, at least 5 bases on each arm are able to bind to, or otherwise interact with, the target RNA.

Example 12: Chemical Synthesis and Purification of Ribozymes and Antisense for Efficient Cleavage and/or Blocking of HER2 Expression

DNAzyme molecules are designed to anneal to various sites in the RNA message. The binding arms of the DNAzyme molecules are complementary to the target site sequences described above. The DNAzymes were chemically synthesized. The method of synthesis used followed the procedure for nucleic acid synthesis as described above and in Usman *et al.*, (1987 *J. Am. Chem. Soc.*, 109, 7845), Scaringe *et al.*, (1990 *Nucleic Acids Res.*, 18, 5433) and Wincott *et al.*, *supra*, and made use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. The average stepwise coupling yields were typically >98%. The sequences of the chemically synthesized DNAzyme molecules used in this study are shown below in Table V.

Example 13: DNAzyme Cleavage of HER2 RNA Target *in vitro*

DNAzymes targeted to the human HER2 RNA are designed and synthesized as described above. These enzymatic nucleic acid molecules can be tested for cleavage activity *in vitro*, for example, using the following procedure. The target sequences and the nucleotide location within the HER2 RNA are given in Tables IV and V.

Cleavage Reactions:

Ribozymes and substrates were synthesized in 96-well format using 0.2 μ mol scale. Substrates were 5'-³²P labeled and gel purified using 7.5% polyacrylamide gels, and eluting into water. Assays were done by combining trace substrate with 500nM Ribozyme or greater, and initiated by adding final concentrations of 40mM Mg⁺², and 50mM Tris-Cl pH 8.0. For each ribozyme/substrate combination a control reaction was done to ensure cleavage was not the result of non-specific substrate degradation. A single three hour time point was taken and run on a 15% polyacrylamide gel to assess cleavage activity. Gels were dried and scanned using a Molecular Dynamics Phosphorimager and quantified using Molecular Dynamics ImageQuant software. Percent cleaved was determined by dividing values for cleaved substrate bands by full-length (uncleaved) values plus cleaved values and multiplying by 100 (%cleaved=[C/(U+C)]*100).

Example 14: DNAzyme Cleavage of HER2 RNA Target *in vivo*

Cell Culture Review

The greatest HER2 specific effects have been observed in cancer cell lines that express high levels of HER2 protein (as measured by ELISA). Specifically, in one study that treated five human breast cancer cell lines with the HER2 antibody (anti-erbB2-sFv), the greatest inhibition of cell growth was seen in three cell lines (MDA-MB-361, SKBR-3 and BT-474) that express high levels of HER2 protein. No inhibition of cell growth was observed in two cell lines (MDA-MB-231 and MCF-7) that express low levels of HER2 protein (Wright, M., Grim, J., Deshane, J., Kim, M., Strong, T.V., Siegel, G.P., Curiel, D.T. (1997) An intracellular anti-erbB-2 single-chain antibody is specifically cytotoxic to human breast carcinoma cells overexpressing erbB-2. *Gene Therapy* 4: 317-322). Another group successfully used SKBR-3 cells to show HER2 antisense oligonucleotide-mediated inhibition of HER2 protein expression and HER2 RNA knockdown (Vaughn, J.P., Iglehart, J.D., Demirdji, S., Davis, P., Babiss, L.E., Caruthers, M.H., Marks, J.R. (1995) Antisense DNA downregulation of the ERBB2 oncogene measured by a flow cytometric assay. *Proc Natl Acad Sci USA* 92: 8338-8342). Other groups have also demonstrated a decrease in the levels of HER2 protein, HER2 mRNA and/or cell proliferation in cultured cells using anti-HER2 DNAzymes or antisense molecules (Suzuki T., Curcio, L.D., Tsai, J. and Kashani-Sabet M. (1997) Anti-c-erb-B-2 Ribozyme for Breast Cancer. In *Methods in Molecular Medicine*, Vol. 11, Therapeutic Applications of Ribozymes, Human Press, Inc., Totowa, NJ; Weichen, K., Zimmer, C. and Dietel, M. (1997) Selection of a high activity c-erbB-2 ribozyme using a

fusion gene of c-erbB-2 and the enhanced green fluorescent protein. *Cancer Gene Therapy* 5: 45-51; Czubayko, F., Downing, S.G., Hsieh, S.S., Goldstein, D.J., Lu P.Y., Trapnell, B.C. and Wellstein, A. (1997) Adenovirus-mediated transduction of ribozymes abrogates HER-2/neu and pleiotrophin expression and inhibits tumor cell proliferation. *Gene Ther.* 4: 943-949; Colomer, R., Lupu, R., Bacus, S.S. and Gelmann, E.P. (1994) *erbB-2* antisense oligonucleotides inhibit the proliferation of breast carcinoma cells with *erbB-2* oncogene amplification. *British J. Cancer* 70: 819-825; Betram *et al.*, 1994). Because cell lines that express higher levels of HER2 have been more sensitive to anti-HER2 agents, we prefer using several medium to high expressing cell lines, including SKBR-3 and T47D, for DNAzyme screens in cell culture.

A variety of endpoints have been used in cell culture models to look at HER2-mediated effects after treatment with anti-HER2 agents. Phenotypic endpoints include inhibition of cell proliferation, apoptosis assays and reduction of HER2 protein expression. Because overexpression of HER2 is directly associated with increased proliferation of breast and ovarian tumor cells, a proliferation endpoint for cell culture assays will preferably be used as the primary screen. There are several methods by which this endpoint can be measured. Following treatment of cells with DNAzymes, cells are allowed to grow (typically 5 days) after which either the cell viability, the incorporation of [³H] thymidine into cellular DNA and/or the cell density can be measured. The assay of cell density is very straightforward and can be done in a 96-well format using commercially available fluorescent nucleic acid stains (such as Syto® 13 or CyQuant®). The assay using CyQuant® is described herein and is currently being employed to screen ~100 DNAzymes targeting HER2 (details below).

As a secondary, confirmatory endpoint a DNAzyme-mediated decrease in the level of HER2 protein expression can be evaluated using a HER2-specific ELISA.

Validation of Cell Lines and DNAzyme Treatment Conditions

Two human breast cancer cell lines (T47D and SKBR-3) that are known to express medium to high levels of HER2 protein, respectively, are considered for DNAzyme screening. In order to validate these cell lines for HER2-mediated sensitivity, both cell lines are treated with the HER2 specific antibody, Herceptin® (Genentech) and its effect on cell proliferation is determined. Herceptin® is added to cells at concentrations ranging from 0–8 µM in medium containing either no serum (OptiMem), 0.1% or 0.5% FBS and efficacy is determined *via* cell proliferation. Maximal inhibition of proliferation (~50%) in both cell lines is typically observed after addition of Herceptin® at 0.5 nM in medium containing 0.1%

or no FBS. The fact that both cell lines are sensitive to an anti-HER2 agent (Herceptin®) supports their use in experiments testing anti-HER2 DNAzymes.

Prior to DNAzyme screening, the choice of the optimal lipid(s) and conditions for DNAzyme delivery is determined empirically for each cell line. Applicant has established a panel of cationic lipids (lipids as described in PCT application WO99/05094) that can be used to deliver DNAzymes to cultured cells and are very useful for cell proliferation assays that are typically 3-5 days in length. (Additional description of useful lipids is provided above, and those skilled in the art are also familiar with a variety of lipids that can be used for delivery of oligonucleotide to cells in culture.) Initially, this panel of lipid delivery vehicles is screened in SKBR-3 and T47D cells using previously established control oligonucleotides. Specific lipids and conditions for optimal delivery are selected for each cell line based on these screens. These conditions are used to deliver HER2 specific DNAzymes to cells for primary (inhibition of cell proliferation) and secondary (decrease in HER2 protein) efficacy endpoints.

Primary Screen: Inhibition of Cell Proliferation

DNAzyme screens are performed using an automated, high throughput 96-well cell proliferation assay. Cell proliferation is measured over a 5-day treatment period using the nucleic acid stain CyQuant® for determining cell density. The growth of cells treated with DNAzyme/lipid complexes is compared to both untreated cells and to cells treated with Scrambled-arm Attenuated core Controls. SACs can no longer bind to the target site due to the scrambled arm sequence and have nucleotide changes in the core that greatly diminish DNAzyme cleavage. These SACs are used to determine non-specific inhibition of cell growth caused by DNAzyme chemistry (*i.e.* multiple 2' O-Me modified nucleotides and a 3' inverted abasic). Lead DNAzymes are chosen from the primary screen based on their ability to inhibit cell proliferation in a specific manner. Dose response assays are carried out on these leads and a subset was advanced into a secondary screen using the level of HER2 protein as an endpoint.

Secondary Screen: Decrease in HER2 Protein and/or RNA

A secondary screen that measures the effect of anti-HER2 DNAzymes on HER2 protein and/or RNA levels is used to affirm preliminary findings. A robust HER2 ELISA for both T47D and SKBR-3 cells has been established and is available for use as an additional endpoint. In addition, a real time RT-PCR assay (TaqMan assay) has been developed to assess HER2 RNA reduction compared to an actin RNA control. Dose response activity of

nucleic acid molecules of the instant invention can be used to assess both HER2 protein and RNA reduction endpoints.

DNAzyme Mechanism Assays

5 A TaqMan® assay for measuring the DNAzyme-mediated decrease in HER2 RNA has also been established. This assay is based on PCR technology and can measure in real time the production of HER2 mRNA relative to a standard cellular mRNA such as GAPDH. This RNA assay is used to establish proof that lead DNAzymes are working through an RNA cleavage mechanism and result in a decrease in the level of HER2 mRNA, thus leading to a decrease in cell surface HER2 protein receptors and a subsequent decrease in tumor cell
10 proliferation.

Animal Models

Evaluating the efficacy of anti-HER2 agents in animal models is an important prerequisite to human clinical trials. As in cell culture models, the most HER2 sensitive mouse tumor xenografts are those derived from human breast carcinoma cells that express
15 high levels of HER2 protein. In a recent study, nude mice bearing BT-474 xenografts were sensitive to the anti-HER2 humanized monoclonal antibody Herceptin®, resulting in an 80% inhibition of tumor growth at a 1 mg kg dose (ip, 2 X week for 4-5 weeks). Tumor eradication was observed in 3 of 8 mice treated in this manner (Baselga, J., Norton, L. Albanell, J., Kim, Y.M. and Mendelsohn, J. (1998) Recombinant humanized anti-HER2
20 antibody (Herceptin) enhances the antitumor activity of paclitaxel and doxorubicin against HER2/neu overexpressing human breast cancer xenografts. *Cancer Res.* 15: 2825-2831). This same study compared the efficacy of Herceptin® alone or in combination with the commonly used chemotherapeutics, paclitaxel or doxorubicin. Although, all three anti-HER2 agents caused modest inhibition of tumor growth, the greatest antitumor activity was
25 produced by the combination of Herceptin® and paclitaxel (93% inhibition of tumor growth vs 35% with paclitaxel alone). The above studies provide proof that inhibition of HER2 expression by anti-HER2 agents causes inhibition of tumor growth in animals. Lead anti-HER2 DNAzymes chosen from *in vitro* assays are further tested in mouse xenograft models. DNAzymes are first tested alone and then in combination with standard chemotherapies.

Animal Model Development

Three human breast tumor cell lines (T47D, SKBR-3 and BT-474) were characterized to establish their growth curves in mice. These three cell lines have been implanted into the

mammary papillae of both nude and SCID mice and primary tumor volumes are measured 3 times per week. Growth characteristics of these tumor lines using a Matrigel implantation format can also be established. The use of two other breast cell lines that have been engineered to express high levels of HER2 can also be used in the described studies. The tumor cell line(s) and implantation method that supports the most consistent and reliable tumor growth is used in animal studies testing the lead HER2 DNAzyme(s). DNAzymes are administered by daily subcutaneous injection or by continuous subcutaneous infusion from Alzet mini osmotic pumps beginning 3 days after tumor implantation and continuing for the duration of the study. Group sizes of at least 10 animals are employed. Efficacy is determined by statistical comparison of tumor volume of DNAzyme-treated animals to a control group of animals treated with saline alone. Because the growth of these tumors is generally slow (45-60 days), an initial endpoint is the time in days it takes to establish an easily measurable primary tumor (i.e. 50-100 mm³) in the presence or absence of DNAzyme treatment.

15 Clinical Summary

Overview

Breast cancer is a common cancer in women and also occurs in men to a lesser degree. The incidence of breast cancer in the United States is ~180,000 cases per year and ~46,000 die each year of the disease. In addition, 21,000 new cases of ovarian cancer per year lead to ~13,000 deaths (data from Hung, M.-C., Matin, A., Zhang, Y., Xing, X., Sorgi, F., Huang, L. and Yu, D. (1995) HER-2/neu-targeting gene therapy - a review. *Gene* 159: 65-71 and the Surveillance, Epidemiology and End Results Program, NCI Surveillance, Epidemiology and End Results Program (SEER) Cancer Statistics Review: http://www.seer.ims.nci.nih.gov/Publications/CSR1973_1996/). Ovarian cancer is a potential secondary indication for anti-HER2 DNAzyme therapy.

A full review of breast cancer is given in the NCI PDQ for Breast Cancer (NCI PDQ/Treatment/Health Professionals/Breast Cancer: http://cancernet.nci.nih.gov/clinpdq/soa/Breast_cancer_Physician.html; NCI PDQ/Treatment/Patients/Breast Cancer: http://cancernet.nci.nih.gov/clinpdq/pif/Breast_cancer_Patient.html). A brief overview is given here. Breast cancer is evaluated or "staged" on the basis of tumor size, and whether it has spread to lymph nodes and/or other parts of the body. In Stage I breast cancer, the cancer

is no larger than 2 centimeters and has not spread outside of the breast. In Stage II, the patient's tumor is 2-5 centimeters but cancer may have spread to the axillary lymph nodes. By Stage III, metastasis to the lymph nodes is typical, and tumors are ≥ 5 centimeters. Additional tissue involvement (skin, chest wall, ribs, muscles *etc.*) may also be noted. Once
5 cancer has spread to additional organs of the body, it is classed as Stage IV.

Almost all breast cancers (>90%) are detected at Stage I or II, but 31% of these are already lymph node positive. The 5-year survival rate for node negative patients (with standard surgery/radiation/chemotherapy /hormone regimens) is 97%; however, involvement of the lymph nodes reduces the 5-year survival to only 77%. Involvement of other organs
10 (\geq Stage III) drastically reduces the overall survival, to 22% at 5 years. Thus, chance of recovery from breast cancer is highly dependent on early detection. Because up to 10% of breast cancers are hereditary, those with a family history are considered to be at high risk for breast cancer and should be monitored very closely.

Therapy

15 Breast cancer is highly treatable and often curable when detected in the early stages. (For a complete review of breast cancer treatments, see the NCI PDQ for Breast Cancer.) Common therapies include surgery, radiation therapy, chemotherapy and hormonal therapy. Depending upon many factors, including the tumor size, lymph node involvement and location of the lesion, surgical removal varies from lumpectomy (removal of the tumor and
20 some surrounding tissue) to mastectomy (removal of the breast, lymph nodes and some or all of the underlying chest muscle). Even with successful surgical resection, as many as 21% of the patients may ultimately relapse (10-20 years). Thus, once local disease is controlled by surgery, adjuvant radiation treatments, chemotherapies and/or hormonal therapies are typically used to reduce the rate of recurrence and improve survival. The therapy regimen
25 employed depends not only on the stage of the cancer at its time of removal, but other variables such the type of cancer (ductal or lobular), whether lymph nodes were involved and removed, age and general health of the patient and if other organs are involved.

Common chemotherapies include various combinations of cytotoxic drugs to kill the cancer cells. These drugs include paclitaxel (Taxol), docetaxel, cisplatin, methotrexate,
30 cyclophosphamide, doxorubin, fluorouracil *etc.* Significant toxicities are associated with these cytotoxic therapies. Well-characterized toxicities include nausea and vomiting,

myelosuppression, alopecia and mucosity. Serious cardiac problems are also associated with certain of the combinations, *e.g.* doxorubin and paclitaxel, but are less common.

Testing for estrogen and progesterone receptors helps to determine whether certain anti-hormone therapies might be helpful in inhibiting tumor growth. If either or both receptors are present, therapies to interfere with the action of the hormone ligands, can be given in combination with chemotherapy and are generally continued for several years. These adjuvant therapies are called SERMs, selective estrogen receptor modulators, and they can give beneficial estrogen-like effects on bone and lipid metabolism while antagonizing estrogen in reproductive tissues. Tamoxifen is one such compound. The primary toxic effect associated with the use of tamoxifen is a 2 to 7-fold increase in the rate of endometrial cancer. Blood clots in the legs and lung and the possibility of stroke are additional side effects. However, tamoxifen has been determined to reduce breast cancer incidence by 49% in high-risk patients and an extensive, somewhat controversial, clinical study is underway to expand the prophylactic use of tamoxifen. Another SERM, raloxifene, was also shown to reduce the incidence of breast cancer in a large clinical trial where it was being used to treat osteoporosis. In additional studies, removal of the ovaries and/or drugs to keep the ovaries from working are being tested.

Bone marrow transplantation is being studied in clinical trials for breast cancers that have become resistant to traditional chemotherapies or where >3 lymph nodes are involved. Marrow is removed from the patient prior to high-dose chemotherapy to protect it from being destroyed, and then replaced after the chemotherapy. Another type of "transplant" involves the exogenous treatment of peripheral blood stem cells with drugs to kill cancer cells prior to replacing the treated cells in the bloodstream.

One biological treatment, a humanized monoclonal anti-HER2 antibody, Herceptin® (Genentech) has been approved by the FDA as an additional treatment for HER2 positive tumors. Herceptin® binds with high affinity to the extracellular domain of HER2 and thus blocks its signaling action. Herceptin® can be used alone or in combination with chemotherapeutics (*i.e.* paclitaxel, docetaxel, cisplatin, *etc.*) (Pegram, M.D., Lipton, A., Hayes, D.F., Weber, B.L., Baselga, J.M., Tripathy, D., Baly, D., Baughman, S.A., Twaddell, T., Glaspy, J.A. and Slamon, D.J. (1998) Phase II study of receptor-enhanced chemosensitivity using recombinant humanized anti-p185HER2/neu monoclonal antibody plus cisplatin in patients with HER2/neu-overexpressing metastatic breast cancer refractory to chemotherapy treatment. *J. Clin. Oncol.* 16: 2659-2671). In Phase III studies, Herceptin® significantly improved the response rate to chemotherapy as well as improving the time to

progression (Ross, J.S. and Fletcher, J.A. (1998) The HER-2/neu oncogene in breast cancer: Prognostic factor, predictive factor and target for therapy. *Oncologist* 3: 1998). The most common side effects attributed to Herceptin® are fever and chills, pain, asthenia, nausea, vomiting, increased cough, diarrhea, headache, dyspnea, infection, rhinitis, and insomnia.

5 Herceptin® in combination with chemotherapy (paclitaxel) can lead to cardiotoxicity (Sparano, J.A. (1999) Doxorubicin/taxane combinations: Cardiac toxicity and pharmacokinetics. *Semin. Oncol.* 26: 14-19), leukopenia, anemia, diarrhea, abdominal pain and infection.

HER2 Protein Levels for Patient Screening and as a Potential Endpoint

- 10 Because elevated HER2 levels can be detected in at least 30% of breast cancers, breast cancer patients can be pre-screened for elevated HER2 prior to admission to initial clinical trials testing an anti-HER2 DNAzyme. Initial HER2 levels can be determined (by ELISA) from tumor biopsies or resected tumor samples.

- During clinical trials, it may be possible to monitor circulating HER2 protein by ELISA
- 15 (Ross and Fletcher, 1998). Evaluation of serial blood/serum samples over the course of the anti-HER2 DNAzyme treatment period could be useful in determining early indications of efficacy. In fact, the clinical course of Stage IV breast cancer was correlated with shed HER2 protein fragment following a dose-intensified paclitaxel monotherapy. In all responders, the HER2 serum level decreased below the detection limit (Luftner, D., Schnabel, S. and
- 20 Possinger, K. (1999) c-erbB-2 in serum of patients receiving fractionated paclitaxel chemotherapy. *Int. J. Biol. Markers* 14: 55-59).

- Two cancer-associated antigens, CA27.29 and CA15.3, can also be measured in the serum. Both of these glycoproteins have been used as diagnostic markers for breast cancer. CA27.29 levels are higher than CA15.3 in breast cancer patients; the reverse is true in healthy
- 25 individuals. Of these two markers, CA27.29 was found to better discriminate primary cancer from healthy subjects. In addition, a statistically significant and direct relationship was shown between CA27.29 and large vs small tumors and node positive vs node negative disease (Gion, M., Mione, R., Leon, A.E. and Dittadi, R. (1999) Comparison of the diagnostic accuracy of CA27.29 and CA15.3 in primary breast cancer. *Clin. Chem.* 45: 630-637).
- 30 Moreover, both cancer antigens were found to be suitable for the detection of possible metastases during follow-up (Rodriguez de Paterna, L., Arnaiz, F., Estenoz, J. Ortuno, B. and Lanzos E. (1999) Study of serum tumor markers CEA, CA15.3, CA27.29 as diagnostic parameters in patients with breast carcinoma. *Int. J. Biol. Markers* 10: 24-29). Thus,

blocking breast tumor growth may be reflected in lower CA27.29 and/or CA15.3 levels compared to a control group. FDA submissions for the use of CA27.29 and CA15.3 for monitoring metastatic breast cancer patients have been filed (reviewed in Beveridge, R.A. (1999) Review of clinical studies of CA27.29 in breast cancer management. *Int. J. Biol. Markers* 14: 36-39). Fully automated methods for measurement of either of these markers are commercially available.

Indications

Particular degenerative and disease states that can be associated with HER2 expression modulation include but are not limited to cancer, for example breast cancer and ovarian cancer and/or any other diseases or conditions that are related to or will respond to the levels of HER2 in a cell or tissue, alone or in combination with other therapies

The present body of knowledge in HER2 research indicates the need for methods to assay HER2 activity and for compounds that can regulate HER2 expression for research, diagnostic, and therapeutic use.

The use of monoclonal antibodies, chemotherapy, radiation therapy, and analgesics, are all non-limiting examples of methods that can be combined with or used in conjunction with the nucleic acid molecules (e.g. DNazymes) of the instant invention. Common chemotherapies that can be combined with nucleic acid molecules of the instant invention include various combinations of cytotoxic drugs to kill cancer cells. These drugs include but are not limited to paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, vinorelbine etc. Those skilled in the art will recognize that other drug compounds and therapies can be similarly be readily combined with the nucleic acid molecules of the instant invention (e.g. DNzyme molecules) are hence within the scope of the instant invention.

Diagnostic uses

The nucleic acid molecules of this invention (e.g., enzymatic nucleic acid molecules) can be used as diagnostic tools to examine genetic drift and mutations within diseased cells or to detect the presence of HER2 RNA in a cell. The close relationship between enzymatic nucleic acid molecule activity and the structure of the target RNA allows the detection of mutations in any region of the molecule that alters the base-pairing and three-dimensional structure of the target RNA. By using multiple enzymatic nucleic acid molecules described in this invention, one can map nucleotide changes which are important to RNA structure and

function *in vitro*, as well as in cells and tissues. Cleavage of target RNAs with enzymatic nucleic acid molecules can be used to inhibit gene expression and define the role (essentially) of specified gene products in the progression of disease. In this manner, other genetic targets can be defined as important mediators of the disease. These experiments can lead to better treatment of the disease progression by affording the possibility of combinational therapies (e.g., multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of enzymatic nucleic acid molecules and/or other chemical or biological molecules). Other *in vitro* uses of enzymatic nucleic acid molecules of this invention are well known in the art, and include detection of the presence of mRNAs associated with HER2-related condition. Such RNA is detected by determining the presence of a cleavage product after treatment with an enzymatic nucleic acid molecule using standard methodology.

In a specific example, enzymatic nucleic acid molecules that cleave only wild-type or mutant forms of the target RNA are used for the assay. The first enzymatic nucleic acid molecule is used to identify wild-type RNA present in the sample and the second enzymatic nucleic acid molecule is used to identify mutant RNA in the sample. As reaction controls, synthetic substrates of both wild-type and mutant RNA are cleaved by both enzymatic nucleic acid molecules to demonstrate the relative enzymatic nucleic acid molecule efficiencies in the reactions and the absence of cleavage of the "non-targeted" RNA species. The cleavage products from the synthetic substrates also serve to generate size markers for the analysis of wild-type and mutant RNAs in the sample population. Thus each analysis requires two enzymatic nucleic acid molecules, two substrates and one unknown sample which is combined into six reactions. The presence of cleavage products is determined using an RNase protection assay so that full-length and cleavage fragments of each RNA can be analyzed in one lane of a polyacrylamide gel. It is not absolutely required to quantify the results to gain insight into the expression of mutant RNAs and putative risk of the desired phenotypic changes in target cells. The expression of mRNA whose protein product is implicated in the development of the phenotype (*i.e.*, HER2) is adequate to establish risk. If probes of comparable specific activity are used for both transcripts, then a qualitative comparison of RNA levels will be adequate and will decrease the cost of the initial diagnosis. Higher mutant form to wild-type ratios are correlated with higher risk whether RNA levels are compared qualitatively or quantitatively. The use of enzymatic nucleic acid molecules in diagnostic applications contemplated by the instant invention is more fully described in George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication

No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, International PCT publication No. WO 99/29842.

Additional Uses

5 Potential uses of sequence-specific enzymatic nucleic acid molecules of the instant invention can have many of the same applications for the study of RNA that DNA restriction endonucleases have for the study of DNA (Nathans *et al.*, 1975 *Ann. Rev. Biochem.* 44:273). For example, the pattern of restriction fragments can be used to establish sequence relationships between two related RNAs, and large RNAs can be specifically cleaved to
10 fragments of a size more useful for study. The ability to engineer sequence specificity of the enzymatic nucleic acid molecule is ideal for cleavage of RNAs of unknown sequence. Applicant has described the use of nucleic acid molecules to modulate gene expression of target genes in bacterial, microbial, fungal, viral, and eukaryotic systems including plant or mammalian cells.

15 All patents and publications mentioned in the specification are indicative of the levels of skill of those skilled in the art to which the invention pertains. All references cited in this disclosure are incorporated by reference to the same extent as if each reference had been incorporated by reference in its entirety individually.

20 One skilled in the art would readily appreciate that the present invention is well adapted to carry out the objects and obtain the ends and advantages mentioned, as well as those inherent therein. The methods and compositions described herein as presently representative of preferred embodiments are exemplary and are not intended as limitations on the scope of the invention. Changes therein and other uses will occur to those skilled in the art, which are encompassed within the spirit of the invention, are defined by the scope of the claims.

25 It will be readily apparent to one skilled in the art that varying substitutions and modifications can be made to the invention disclosed herein without departing from the scope and spirit of the invention. Thus, such additional embodiments are within the scope of the present invention and the following claims.

30 The invention illustratively described herein suitably can be practiced in the absence of any element or elements, limitation or limitations which is not specifically disclosed herein. Thus, for example, in each instance herein any of the terms "comprising", "consisting

essentially of' and "consisting of" can be replaced with either of the other two terms. The terms and expressions that have been employed are used as terms of description and not of limitation, and there is no intention that in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof, but it is recognized
5 that various modifications are possible within the scope of the invention claimed. Thus, it should be understood that although the present invention has been specifically disclosed by preferred embodiments, optional features, modification and variation of the concepts herein disclosed can be resorted to by those skilled in the art, and that such modifications and variations are considered to be within the scope of this invention as defined by the description
10 and the appended claims.

In addition, where features or aspects of the invention are described in terms of Markush groups or other grouping of alternatives, those skilled in the art will recognize that the invention is also thereby described in terms of any individual member or subgroup of members of the Markush group or other group.

15 Other embodiments are within the claims that follow.

Table I:**A. 2.5 μ mol Synthesis Cycle ABI 394 Instrument**

Reagent	Equivalents	Amount	Wait Time* DNA	Wait Time* 2'-O-methyl	Wait Time*RNA
Phosphoramidites	6.5	163 μ L	45 sec	2.5 min	7.5 min
S-Ethyl Tetrazole	23.8	238 μ L	45 sec	2.5 min	7.5 min
Acetic Anhydride	100	233 μ L	5 sec	5 sec	5 sec
N-Methyl Imidazole	186	233 μ L	5 sec	5 sec	5 sec
TCA	176	2.3 mL	21 sec	21 sec	21 sec
Iodine	11.2	1.7 mL	45 sec	45 sec	45 sec
Beaucage	12.9	645 μ L	100 sec	300 sec	300 sec
Acetonitrile	NA	6.67 mL	NA	NA	NA

B. 0.2 μ mol Synthesis Cycle ABI 394 Instrument

Reagent	Equivalents	Amount	Wait Time* DNA	Wait Time* 2'-O-methyl	Wait Time*RNA
Phosphoramidites	15	31 μ L	45 sec	233 sec	465 sec
S-Ethyl Tetrazole	38.7	31 μ L	45 sec	233 min	465 sec
Acetic Anhydride	655	124 μ L	5 sec	5 sec	5 sec
N-Methyl Imidazole	1245	124 μ L	5 sec	5 sec	5 sec
TCA	700	732 μ L	10 sec	10 sec	10 sec
Iodine	20.6	244 μ L	15 sec	15 sec	15 sec
Beaucage	7.7	232 μ L	100 sec	300 sec	300 sec
Acetonitrile	NA	2.64 mL	NA	NA	NA

C. 0.2 μ mol Synthesis Cycle 96 well Instrument

Reagent	Equivalents:DNA/ 2'-O-methyl/Ribo	Amount: DNA/2'-O- methyl/Ribo	Wait Time* DNA	Wait Time* 2'-O- methyl	Wait Time* Ribo
Phosphoramidites	22/33/66	40/60/120 μ L	60 sec	180 sec	360sec
S-Ethyl Tetrazole	70/105/210	40/60/120 μ L	60 sec	180 min	360 sec
Acetic Anhydride	265/265/265	50/50/50 μ L	10 sec	10 sec	10 sec
N-Methyl Imidazole	502/502/502	50/50/50 μ L	10 sec	10 sec	10 sec
TCA	238/475/475	250/500/500 μ L	15 sec	15 sec	15 sec
Iodine	6.8/6.8/6.8	80/80/80 μ L	30 sec	30 sec	30 sec
Beaucage	34/51/51	80/120/120	100 sec	200 sec	200 sec
Acetonitrile	NA	1150/1150/1150 μ L	NA	NA	NA

- Wait time does not include contact time during delivery.

Table II: Human K-Ras DNzyme and Substrate Sequence

Pos	Substrate	Seq ID	DNzyme	Seq ID
10	CCUAGGCG G CGGCCGCG	1	CGCGGCCG GGCTAGCTACAACGA CGCCTAGG	2329
13	AGGCGGCG G CCGCGGCG	2	CGCCGCGG GGCTAGCTACAACGA CGCCGCCT	2330
16	CGGCGGCC G CGGCGGCG	3	CGCCGCGG GGCTAGCTACAACGA GGCCGCCG	2331
19	CGGCCGCG G CGGCGGAG	4	CTCCGCGG GGCTAGCTACAACGA CGCGGCCG	2332
22	CCGCGGCG G CGGAGGCA	5	TGCCTCCG GGCTAGCTACAACGA CGCCGCGG	2333
28	CGGCGGAG G CAGCAGCG	6	CGCTGCTG GGCTAGCTACAACGA CTCCGCGG	2334
31	CGGAGGCA G CAGCGGCG	7	CGCCGCTG GGCTAGCTACAACGA TGCCTCCG	2335
34	AGGCAGCA G CGGCGGCG	8	CGCCGCGG GGCTAGCTACAACGA TGCTGCCT	2336
37	CAGCAGCG G CGGCGGCA	9	TGCCGCGG GGCTAGCTACAACGA CGCTGCTG	2337
40	CAGCGGCG G CGGCAGUG	10	CACTGCCG GGCTAGCTACAACGA CGCCGCTG	2338
43	CGGCGGCG G CAGUGGCG	11	CGCCACTG GGCTAGCTACAACGA CGCCGCCG	2339
46	CGGCGGCA G UGGCGGCG	12	CGCCGCCA GGCTAGCTACAACGA TGCCGCCG	2340
49	CGGCAGUG G CGGCGGCG	13	CGCCGCCG GGCTAGCTACAACGA CACTGCCG	2341
52	CAGUGGCG G CGGCGAAG	14	CTTCGCCG GGCTAGCTACAACGA CGCCACTG	2342
55	UGGCGGCG G CGAAGGUG	15	CACCTTCG GGCTAGCTACAACGA CGCCGCCA	2343
61	CGGCGAAG G UGGCGGCG	16	CGCCGCCA GGCTAGCTACAACGA CTTCGCCG	2344
64	CGAAGGUG G CGGCGGCU	17	AGCCGCCG GGCTAGCTACAACGA CACCTTCG	2345
67	AGGUGGCG G CGGCUCGG	18	CCGAGCCG GGCTAGCTACAACGA CGCCACCT	2346
70	UGGCGGCG G CUCGGCCA	19	TGGCCGAG GGCTAGCTACAACGA CGCCGCCA	2347
75	GCGGCUCG G CCAGUACU	20	AGTACTGG GGCTAGCTACAACGA CGAGCCGC	2348
79	CUCGGCCA G UACUCCCG	21	CGGGAGTA GGCTAGCTACAACGA TGGCCGAG	2349
81	CGGCCAGU A CUCCCGGC	22	GCCGGGAG GGCTAGCTACAACGA ACTGGCCG	2350
88	UACUCCCG G CCCC CGCC	23	GGCGGGGG GGCTAGCTACAACGA CGGGAGTA	2351
94	CGGCCCCC G CCAUUUCG	24	CGAAATGG GGCTAGCTACAACGA GGGGGCCG	2352
97	CCCCCGCC A UUUCGGAC	25	GTCCGAAA GGCTAGCTACAACGA GGCGGGGG	2353
104	CAUUUCGG A CUGGGAGC	26	GCTCCAG GGCTAGCTACAACGA CCGAAATG	2354
111	GACUGGGA G CGAGCGCG	27	CGCGCTCG GGCTAGCTACAACGA TCCAGTC	2355
115	GGGAGCGA G CGCGCGCG	28	GCGCCGCG GGCTAGCTACAACGA TCGTCCC	2356
117	GAGCGAGC G CGGCGCAG	29	CTGCGCCG GGCTAGCTACAACGA GCTCGCTC	2357
120	CGAGCGCG G CGCAGGCA	30	TGCCTGCG GGCTAGCTACAACGA CGCGCTCG	2358
122	AGCGCGGC G CAGGCACU	31	AGTGCCTG GGCTAGCTACAACGA GCCGCGCT	2359
126	CGGCGCAG G CACUGAAG	32	CTTCAGTG GGCTAGCTACAACGA CTGCGCCG	2360
128	GCGCAGGC A CUGAAGGC	33	GCCTTCAG GGCTAGCTACAACGA GCCTGCGC	2361
135	CACUGAAG G CGGCGGCG	34	CGCCGCCG GGCTAGCTACAACGA CTTCAGTG	2362
138	UGAAGGCG G CGGCGGGG	35	CCCCGCCG GGCTAGCTACAACGA CGCCTTCA	2363
141	AGGCGGCG G CGGGGCCA	36	TGGCCCCG GGCTAGCTACAACGA CGCCGCCT	2364
146	GCGCGGGG G CCAGAGGC	37	GCCTCTGG GGCTAGCTACAACGA CCCGCCGC	2365
153	GGCCAGAG G CUCAGCGG	38	CCGCTGAG GGCTAGCTACAACGA CTCTGGCC	2366
158	GAGGCUCA G CGGCUCCC	39	GGGAGCCG GGCTAGCTACAACGA TGAGCCTC	2367
161	GCUCAGCG G CUCCAGG	40	CCTGGGAG GGCTAGCTACAACGA CGCTGAGC	2368
169	GCUCCAG G UGCGGGAG	41	CTCCCGCA GGCTAGCTACAACGA CTGGGAGC	2369
171	UCCAGGU G CGGGAGAG	42	CTCTCCCG GGCTAGCTACAACGA ACCTGGGA	2370
182	GGAGAGAG G CCUGCUGA	43	TCAGCAGG GGCTAGCTACAACGA CTCTCTCC	2371
186	AGAGCCCU G CUGAAAAU	44	ATTTTCAG GGCTAGCTACAACGA AGGCCTCT	2372
193	UGCUGAAA A UGACUGAA	45	TTCAGTCA GGCTAGCTACAACGA TTTCAGCA	2373
196	UGAAAAUG A CUGAAUUAU	46	ATATTTCAG GGCTAGCTACAACGA CATTTTCA	2374
201	AUGACUGA A UAUAAACU	47	AGTTTATA GGCTAGCTACAACGA TCAGTCAT	2375
203	GACUGAAU A UAAACUUG	48	CAAGTTTA GGCTAGCTACAACGA ATTTCAGT	2376

207	GAAUAUAA A CUUGUGGU	49	ACCACAAG GGCTAGCTACAACGA TTATATTC	2377
211	AUAAACUU G UGGUAGUU	50	AACTACCA GGCTAGCTACAACGA AAGTTTAT	2378
214	AACUUGUG G UAGUUGGA	51	TCCAACCTA GGCTAGCTACAACGA CACAAGTT	2379
217	UUGUGGUA G UUGGAGCU	52	AGCTCCAA GGCTAGCTACAACGA TACCACAA	2380
223	UAGUUGGA G CUUGUGGC	53	GCCACAAG GGCTAGCTACAACGA TCCAACCTA	2381
227	UGGAGCUU G UGGCGUAG	54	CTACGCCA GGCTAGCTACAACGA AAGCTCCA	2382
230	AGCUUGUG G CGUAGGCA	55	TGCCTACG GGCTAGCTACAACGA CACAAGCT	2383
232	CUUGUGGC G UAGGCAAG	56	CTTGCCTA GGCTAGCTACAACGA GCCACAAG	2384
236	UGGCGUAG G CAAGAGUG	57	CACTCTTG GGCTAGCTACAACGA CTACGCCA	2385
242	AGGCAAGA G UGCCUUGA	58	TCAAGGCA GGCTAGCTACAACGA TCTTGCCT	2386
244	GCAAGAGU G CCUUGACG	59	CGTCAAGG GGCTAGCTACAACGA ACTCTTGC	2387
250	GUGCCUUG A CGAUACAG	60	CTGTATCG GGCTAGCTACAACGA CAAGGCAC	2388
253	CCUUGACG A UACAGCUA	61	TAGCTGTA GGCTAGCTACAACGA CGTCAAGG	2389
255	UUGACGAU A CAGCUAAU	62	ATTAGCTG GGCTAGCTACAACGA ATCGTCAA	2390
258	ACGAUACA G CUAUUAU	63	TGAATTAG GGCTAGCTACAACGA TGTATCGT	2391
262	UACAGCUA A UUCAGAAU	64	ATTCTGAA GGCTAGCTACAACGA TAGCTGTA	2392
269	AAUUCAGA A UCAUUUUG	65	CAAAATGA GGCTAGCTACAACGA TCTGAATT	2393
272	UCAGAAUC A UUUUGUGG	66	CCACAAAA GGCTAGCTACAACGA GATTCTGA	2394
277	AUCAUUUU G UGGACGAA	67	TTCGTCCA GGCTAGCTACAACGA AAAATGAT	2395
281	UUUUGUGG A CGAAUAUG	68	CATATTCG GGCTAGCTACAACGA CCACAAAA	2396
285	GUGGACGA A UAUGAUCC	69	GGATCATA GGCTAGCTACAACGA TCGTCCAC	2397
287	GGACGAU A UGAUCCAA	70	TTGGATCA GGCTAGCTACAACGA ATTCTGCC	2398
290	CGAAUAUG A UCCAACAA	71	TTGTTGGA GGCTAGCTACAACGA CATATTCG	2399
295	AUGAUCCA A CAUAGAG	72	CTCTATTG GGCTAGCTACAACGA TGGATCAT	2400
298	AUCCAACA A UAGAGGAU	73	ATCCTCTA GGCTAGCTACAACGA TGTGGAT	2401
305	AAUAGAGG A UUCUACA	74	TGTAGGAA GGCTAGCTACAACGA CCTCTATT	2402
311	GGAUUCCU A CAGGAAGC	75	GCTTCCTG GGCTAGCTACAACGA AGGAATCC	2403
318	UACAGGAA G CAAGUAGU	76	ACTACTTG GGCTAGCTACAACGA TTCTGTGA	2404
322	GGAAAGCA G UAGUAAU	77	AATTACTA GGCTAGCTACAACGA TTGCTTCC	2405
325	AGCAAGUA G UAAUUGAU	78	ATCAATTA GGCTAGCTACAACGA TACTTGCT	2406
328	AAGUAGUA A UGAUGGA	79	TCCATCAA GGCTAGCTACAACGA TACTACTT	2407
332	AGUAAUUG A UGGAGAAA	80	TTTCTCCA GGCTAGCTACAACGA CAATTACT	2408
340	AUGGAGAA A CCUGUCUC	81	GAGACAGG GGCTAGCTACAACGA TTCTCCAT	2409
344	AGAAACCU G UCUCUUGG	82	CCAAGAGA GGCTAGCTACAACGA AGGTTTCT	2410
353	UCUCUUGG A UAUUCUCG	83	CGAGAATA GGCTAGCTACAACGA CCAAGAGA	2411
355	UCUUGGAU A UUCUCGAC	84	GTCGAGAA GGCTAGCTACAACGA ATCCAAGA	2412
362	UAUUCUCG A CACAGCAG	85	CTGCTGTG GGCTAGCTACAACGA CGAGAATA	2413
364	UUCUCGAC A CAGCAGGU	86	ACCTGCTG GGCTAGCTACAACGA GTCGAGAA	2414
367	UCGACACA G CAGGUCAA	87	TTGACCTG GGCTAGCTACAACGA TGTGTCGA	2415
371	CACAGCAG G UCAAGAGG	88	CCTCTTGA GGCTAGCTACAACGA CTGCTGTG	2416
381	CAAGAGGA G UACAGUGC	89	GCACTGTA GGCTAGCTACAACGA TCCTCTTG	2417
383	AGAGGAGU A CAGUGCAA	90	TTGCACTG GGCTAGCTACAACGA ACTCCTCT	2418
386	GGAGUACA G UGCAUAGA	91	TCATTGCA GGCTAGCTACAACGA TGTACTCC	2419
388	AGUACAGU G CAAUGAGG	92	CCTCATTG GGCTAGCTACAACGA ACTGTACT	2420
391	ACAGUGCA A UGAGGGAC	93	GTCCCTCA GGCTAGCTACAACGA TGCACTGT	2421
398	AAUGAGGG A CCAGUACA	94	TGTACTGG GGCTAGCTACAACGA CCCTCATT	2422
402	AGGGACCA G UACAUGAG	95	CTCATGTA GGCTAGCTACAACGA TGGTCCCT	2423
404	GGACCAGU A CAUGAGGA	96	TCCTCATG GGCTAGCTACAACGA ACTGGTCC	2424
406	ACCAGUAC A UGAGGACU	97	AGTCTCA GGCTAGCTACAACGA GTACTGGT	2425
412	ACAUGAGG A CUGGGGAG	98	CTCCCCAG GGCTAGCTACAACGA CCTCATGT	2426
422	UGGGGAGG G CUUUCUUU	99	AAAGAAAG GGCTAGCTACAACGA CCTCCCCA	2427
431	CUUUCUUU G UGUUUUG	100	CAAATACA GGCTAGCTACAACGA AAAGAAAG	2428

433	UUCUUUGU G UAUUUGCC	101	GGCAAATA GGCTAGCTACAACGA ACAAAGAA	2429
435	CUUUGUGU A UUUGCCAU	102	ATGGCAAA GGCTAGCTACAACGA ACACAAAG	2430
439	GUGUAUUU G CCAUAAAU	103	ATTTATGG GGCTAGCTACAACGA AAATACAC	2431
442	UAUUUGCC A UAAUAAU	104	ATTATTTA GGCTAGCTACAACGA GGCAAATA	2432
446	UGCCAUA A UAAUACUA	105	TAGTATTA GGCTAGCTACAACGA TTATGGCA	2433
449	CAUAAUA A UACUAAU	106	ATTTAGTA GGCTAGCTACAACGA TATTTATG	2434
451	UAAUAAU A CUAAUCA	107	TGATTTAG GGCTAGCTACAACGA ATTATTTA	2435
456	AAUACUA A UCAUUUGA	108	TCAAATGA GGCTAGCTACAACGA TTAGTATT	2436
459	ACUAAUC A UUUGAAGA	109	TCTTCAA GGCTAGCTACAACGA GATTTAGT	2437
467	AUUUGAAG A UAUUCACC	110	GGTGAATA GGCTAGCTACAACGA CTTCAAAT	2438
469	UUGAAGAU A UUCACCAU	111	ATGGTGAA GGCTAGCTACAACGA ATCTTCAA	2439
473	AGAUUUC A CCAUUAU	112	TATAATGG GGCTAGCTACAACGA GAATATCT	2440
476	UAUUCACC A UUAUAGAG	113	CTCTATAA GGCTAGCTACAACGA GGTGAATA	2441
479	UCACCAU A UAGAGAAC	114	GTTCTCTA GGCTAGCTACAACGA AATGGTGA	2442
486	UAUAGAGA A CAAAUUA	115	TTAATTTG GGCTAGCTACAACGA TCTCTATA	2443
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517	CUGAAGAU G UACCUAUG	120	CATAGGTA GGCTAGCTACAACGA ATCTTCAG	2448
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526	UACCUAUG G UCCUAGUA	123	TACTAGGA GGCTAGCTACAACGA CATAGGTA	2451
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539	AGUAGGAA A UAAUUGUG	125	CACATTTA GGCTAGCTACAACGA TTCCTACT	2453
543	GGAAUUA A UGUGAUUU	126	AAATCACA GGCTAGCTACAACGA TTATTTCC	2454
545	AAAUAAAU G UGAUUUGC	127	GCAAATCA GGCTAGCTACAACGA ATTTATTT	2455
548	UAAUUGUG A UUUGCCUU	128	AAGGCAAA GGCTAGCTACAACGA CACATTTA	2456
552	UGUGAUUU G CCUUCUAG	129	CTAGAAGG GGCTAGCTACAACGA AAATCACA	2457
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580	CAAAACAG G CUCAGGAC	135	GTCCTGAG GGCTAGCTACAACGA CTGTTTTG	2463
587	GGCUCAGG A CUUAGCAA	136	TTGCTAAG GGCTAGCTACAACGA CCTGAGCC	2464
592	AGGACUUA G CAAGAAGU	137	ACTTCTTG GGCTAGCTACAACGA TAAGTCCT	2465
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644	AAGACAGG G UGUUGAUG	147	CATCAACA GGCTAGCTACAACGA CCTGTCTT	2475
646	GACAGGGU G UUGAUGAU	148	ATCATCAA GGCTAGCTACAACGA ACCCTGTC	2476
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655	UUGAUGAU G CCUUCUUA	151	ATAGAAGG GGCTAGCTACAACGA ATCATCAA	2479
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670	AUACAUUA G UUCGAGAA	155	TTCTCGAA GGCTAGCTACAACGA TAATGTAT	2483
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799	UAGUACAA G UGUAAUU	181	AATTACCA GGCTAGCTACAACGA TTGTACTA	2509
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820	UACAUUAC A CUAAUUA	188	TAATTTAG GGCTAGCTACAACGA GTAATGTA	2516
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834	UUAUAGC A UUUGUUU	192	AAAACAAA GGCTAGCTACAACGA GCTAATAA	2520
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844	UUGUUUUA G CAUUACCU	194	AGGTAATG GGCTAGCTACAACGA TAAAACAA	2522
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922	CAGUGGAA G UUUUUUUU	211	AAAAAAA GGCTAGCTACAACGA TTCCACTG	2539
939	UCCUCGAA G UGCCAGUA	212	TACTGGCA GGCTAGCTACAACGA TTCGAGGA	2540
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1530	AUAUUUAC A UGCUACUA	342	TAGTAGCA GGCTAGCTACAACGA GTAAATAT	2670
1532	AUUUACAU G CUACUAAA	343	TTTAGTAG GGCTAGCTACAACGA ATGTAAAT	2671
1535	UACAUGC A CUAAUUU	344	AAATTTAG GGCTAGCTACAACGA AGCATGTA	2672
1540	GCUCUAA A UUUUUAUA	345	TATAAAAA GGCTAGCTACAACGA TTAGTAGC	2673
1546	AAAUUUUU A UAAUAAU	346	AATTATTA GGCTAGCTACAACGA AAAAATTT	2674
1549	UUUUUAUA A UAAUUGAA	347	TTCAATTA GGCTAGCTACAACGA TATAAAAA	2675
1552	UUUAUAUA A UUGAAAAG	348	CTTTTCAA GGCTAGCTACAACGA TATTATAA	2676
1561	UUGAAAAG A UUUUAACA	349	TGTTAAAA GGCTAGCTACAACGA CTTTTC	2677
1567	AGAUUUUA A CAAGUAUA	350	TATACTTG GGCTAGCTACAACGA TAAAATCT	2678
1571	UUUAACAA G UAUAAAA	351	TTTTTATA GGCTAGCTACAACGA TTGTTAAA	2679
1573	UAACAAGU A UAAAAAA	352	TTTTTTTA GGCTAGCTACAACGA ACTTGTTA	2680
1581	AUAAAAA A UUCUCAUA	353	TATGAGAA GGCTAGCTACAACGA TTTTTTAT	2681
1587	AAAUUCUC A UAGGAUU	354	AATTCCTA GGCTAGCTACAACGA GAGAATTT	2682
1593	UCAUAGGA A UUAUUGU	355	ACATTTAA GGCTAGCTACAACGA TCCTATGA	2683
1598	GGAAUUA A UGUAGUCU	356	AGACTACA GGCTAGCTACAACGA TTAATTCC	2684
1600	AAUUAAAU G UAGUCUCC	357	GGAGACTA GGCTAGCTACAACGA ATTTAATT	2685
1603	UAAUUGUA G UCUCUCC	358	CAGGGAGA GGCTAGCTACAACGA TACATTTA	2686
1611	GUCUCCCU G UGUCAGAC	359	GTCTGACA GGCTAGCTACAACGA AGGGAGAC	2687
1613	CUCCUGU G UCAGACUG	360	CAGTCTGA GGCTAGCTACAACGA ACAGGGAG	2688

1618	UGUGUCAG A CUGCUCUU	361	AAGAGCAG GGCTAGCTACAACGA CTGACACA	2689
1621	GUCAGACU G CUCUUUCA	362	TGAAAGAG GGCTAGCTACAACGA AGTCTGAC	2690
1629	GCUCUUUC A UAGUAUAA	363	TTATACTA GGCTAGCTACAACGA GAAAGAGC	2691
1632	CUUUCAUA G UUAACUUU	364	AAGTTATA GGCTAGCTACAACGA TATGAAAG	2692
1634	UUCAUAGU A UAACUUUA	365	TAAAGTTA GGCTAGCTACAACGA ACTATGAA	2693
1637	AUAGUAUA A CUUUAUUU	366	ATTTAAAG GGCTAGCTACAACGA TATACTAT	2694
1644	AACUUUAA A UCUUUUCU	367	AGAAAAGA GGCTAGCTACAACGA TTAAAGTT	2695
1656	UUUCUUCA A CUUGAGUC	368	GACTCAAG GGCTAGCTACAACGA TGAAGAAA	2696
1662	CAACUUGA G UCUUGAGG	369	TTCAAAGA GGCTAGCTACAACGA TCAAGTTG	2697
1672	CUUGAAG A UAGUUUUA	370	TAAACTA GGCTAGCTACAACGA CTTCAAAG	2698
1675	UGAAGUA G UUUUAAU	371	AATTAAAA GGCTAGCTACAACGA TATCTTCA	2699
1681	UAGUUUUA A UUCUGCUU	372	AAGCAGAA GGCTAGCTACAACGA TAAACTA	2700
1686	UUAUUUCU G CUUGUGAC	373	GTCACAAG GGCTAGCTACAACGA AGAATTAA	2701
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1693	UGCUGUG A CAUUAUAA	375	TTTTAATG GGCTAGCTACAACGA CACAAGCA	2703
1695	CUUGUGAC A UUAUAAAG	376	TCTTTTAA GGCTAGCTACAACGA GTCACAAG	2704
1703	AUUAUAAAG A UUAUUUGG	377	CCAAATAA GGCTAGCTACAACGA CTTTAAAT	2705
1706	AAAAGAUU A UUUGGGCC	378	GGCCCCAA GGCTAGCTACAACGA AATCTTTT	2706
1712	UUAUUUGG G CCAGUUUAU	379	ATAACTGG GGCTAGCTACAACGA CCAAATAA	2707
1716	UUUGGGCA G UUAUAGCU	380	AGCTATAA GGCTAGCTACAACGA TGGCCCCA	2708
1719	GGCCAGUU A UAGCUUAU	381	ATAAGCTA GGCTAGCTACAACGA AACTGGCC	2709
1722	CAGUUUAU G CUUAUUAG	382	CTAATAAG GGCTAGCTACAACGA TATAACTG	2710
1726	UAUAGCUU A UUAGGUGU	383	ACACCTAA GGCTAGCTACAACGA AAGCTATA	2711
1731	CUUAUUAG G UGUUGAAG	384	CTTCAACA GGCTAGCTACAACGA CTAATAAG	2712
1733	UAUUAGGU G UUGAAGAG	385	CTCTTCAA GGCTAGCTACAACGA ACCTAATA	2713
1742	UUGAAGAG A CCAAGGUU	386	AACCTTGG GGCTAGCTACAACGA CTCTTCAA	2714
1748	AGACCAAG G UUGCAAGC	387	GCTTGCAA GGCTAGCTACAACGA CTTGGTCT	2715
1751	CCAAGGUU G CAAGCCAG	388	CTGGCTTG GGCTAGCTACAACGA AACCTTGG	2716
1755	GGUUGCAA G CCAGGCCC	389	GGGCCTGG GGCTAGCTACAACGA TTGCAACC	2717
1760	CAAGCCAG G CCCUGUGU	390	ACACAGGG GGCTAGCTACAACGA CTGGCTTG	2718
1765	CAGGCCCU G UGUGAACC	391	GGTTCACA GGCTAGCTACAACGA AGGGCCTG	2719
1767	GGCCUGU G UGAACCUU	392	AAGGTTCA GGCTAGCTACAACGA ACAGGGCC	2720
1771	CUGUGUGA A CCUGAGC	393	GCTCAAGG GGCTAGCTACAACGA TCACACAG	2721
1778	AACCUUGA G CUUUCAUA	394	TATGAAAG GGCTAGCTACAACGA TCAAGGTT	2722
1784	GAGCUUUC A UAGAGAGU	395	ACTCTCTA GGCTAGCTACAACGA GAAAGCTC	2723
1791	CAUAGAGA G UUUCACAG	396	CTGTGAAA GGCTAGCTACAACGA TCTCTATG	2724
1796	AGAGUUUC A CAGCAUGG	397	CCATGCTG GGCTAGCTACAACGA GAAACTCT	2725
1799	GUUUCACA G CAUGGACU	398	AGTCCATG GGCTAGCTACAACGA TGTGAAAC	2726
1801	UUCACAGC A UGGACUGU	399	ACAGTCCA GGCTAGCTACAACGA GCTGTGAA	2727
1805	CAGCAUGG A CUGUGUGC	400	GCACACAG GGCTAGCTACAACGA CCATGCTG	2728
1808	CAUGGACU G UGUGCCCC	401	GGGGCACA GGCTAGCTACAACGA AGTCCATG	2729
1810	UGGACUGU G UGCCCCAC	402	GTGGGGCA GGCTAGCTACAACGA ACAGTCCA	2730
1812	GACUGUGU G CCCCACGG	403	CCGTGGGG GGCTAGCTACAACGA ACACAGTC	2731
1817	UGUGCCCC A CGGUCAUC	404	GATGACCG GGCTAGCTACAACGA GGGGCACA	2732
1820	GCCCCACG G UCAUCCGA	405	TCGGATGA GGCTAGCTACAACGA CGTGGGGC	2733
1823	CCACGGUC A UCCGAGUG	406	CACTCGGA GGCTAGCTACAACGA GACCGTGG	2734
1829	UCAUCCGA G UGGUUGUA	407	TACAACCA GGCTAGCTACAACGA TCGGATGA	2735
1832	UCCGAGUG G UUGUACGA	408	TCGTACAA GGCTAGCTACAACGA CACTCGGA	2736
1835	GAGUGGUU G UACGAUGC	409	GCATCGTA GGCTAGCTACAACGA AACCCTC	2737
1837	GUGGUUGU A CGAUGCAU	410	ATGCATCG GGCTAGCTACAACGA ACAACCAC	2738
1840	GUUGUACG A UGCAUUGG	411	CCAATGCA GGCTAGCTACAACGA CGTACAAC	2739
1842	UGUACGAU G CAUUGGUU	412	AACCAATG GGCTAGCTACAACGA ATCGTACA	2740

1844	UACGAUGC A UUGGUUAG	413	CTAACCAA GGCTAGCTACAACGA GCATCGTA	2741
1848	AUGCAUUG G UUAGUCAA	414	TTGACTAA GGCTAGCTACAACGA CAATGCAT	2742
1852	AUUGGUUA G UCAAAAAU	415	ATTTTGA GGCTAGCTACAACGA TAACCAAT	2743
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1869	GGGGAGGG A CUAGGGCA	417	TGCCCTAG GGCTAGCTACAACGA CCCTCCCC	2745
1875	GGACUAGG G CAGUUUGG	418	CCAACTG GGCTAGCTACAACGA CCTAGTCC	2746
1878	CUAGGGCA G UUUGGAUA	419	TATCCAAA GGCTAGCTACAACGA TGCCCTAG	2747
1884	CAGUUUGG A UAGCUCAA	420	TTGAGCTA GGCTAGCTACAACGA CCAAACTG	2748
1887	UUUGGAUA G CUCAACAA	421	TTGTTGAG GGCTAGCTACAACGA TATCCAAA	2749
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1897	UCAACAAG A UACAAUCU	423	AGATTGTA GGCTAGCTACAACGA CTTGTTGA	2751
1899	AACAAGAU A CAAUCUCA	424	TGAGATTG GGCTAGCTACAACGA ATCTTGTT	2752
1902	AAGAUACA A UCUCACUC	425	GAGTGAGA GGCTAGCTACAACGA TGTATCTT	2753
1907	ACAAUCUC A CUCUGUGG	426	CCACAGAG GGCTAGCTACAACGA GAGATTGT	2754
1912	CUCACUCU G UGGUGGUC	427	GACCACCA GGCTAGCTACAACGA AGAGTGAG	2755
1915	ACUCUGUG G UGGUCCUG	428	CAGGACCA GGCTAGCTACAACGA CACAGAGT	2756
1918	CUGUGGUG G UCCUGCUG	429	CAGCAGGA GGCTAGCTACAACGA CACCACAG	2757
1923	GUGGUCCU G CUGACAAA	430	TTTGTCAG GGCTAGCTACAACGA AGGACCAC	2758
1927	UCCUGCUG A CAAAUCAA	431	TTGATTG GGCTAGCTACAACGA CAGCAGGA	2759
1931	GCUGACAA A UCAAGAGC	432	GCTCTTGA GGCTAGCTACAACGA TTGTCAGC	2760
1938	AAUCAAGA G CAUUGCUU	433	AAGCAATG GGCTAGCTACAACGA TCTTGATT	2761
1940	UCAAGAGC A UUGCUUUU	434	AAAAGCAA GGCTAGCTACAACGA GCTCTTGA	2762
1943	AGAGCAUU G CUUUUGUU	435	AACAAAAG GGCTAGCTACAACGA AATGCTCT	2763
1949	UUGCUUUU G UUUCUUA	436	TTAAGAAA GGCTAGCTACAACGA AAAAGCAA	2764
1962	UUAAGAAA A CAAACUCU	437	AGAGTTTG GGCTAGCTACAACGA TTTCTTAA	2765
1966	GAAAACAA A CUCUUUUU	438	AAAAGAG GGCTAGCTACAACGA TTGTTTTC	2766
1980	UUUUAAAA A UUACUUUU	439	AAAAGTAA GGCTAGCTACAACGA TTTTAAAA	2767
1983	UAAAAAUU A CUUUUAAA	440	TTTAAAA GGCTAGCTACAACGA AATTTTAA	2768
1991	ACUUUUAA A UAUUAACU	441	AGTTAATA GGCTAGCTACAACGA TTAAGT	2769
1993	UUUUAAAU A UUAACUCA	442	TGAGTTAA GGCTAGCTACAACGA ATTTAAAA	2770
1997	AAAUUAUA A CUCAAAAG	443	CTTTTGAG GGCTAGCTACAACGA TAATATT	2771
2005	ACUCAAAA G UUGAGAUU	444	AATCTCAA GGCTAGCTACAACGA TTTTGAGT	2772
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2027	GUGGUGGU G UGCCAAGA	449	TCTTGGCA GGCTAGCTACAACGA ACCACCAC	2777
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2035	GUGCCAAG A CAUUAUUU	451	AATTAATG GGCTAGCTACAACGA CTGGGCAC	2779
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2054	UUUUUUA A CAUGAAG	454	CTTCATTG GGCTAGCTACAACGA TTAAAAAA	2782
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2062	ACAAUGAA G UGAAAAAG	456	CTTTTCA GGCTAGCTACAACGA TTCATTGT	2784
2070	GUGAAAAA G UUUUACAA	457	TTGTAAAA GGCTAGCTACAACGA TTTTTCAC	2785
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2564	CUUCUUCU A UAUUAGUG	567	CACTAATA GGCTAGCTACAACGA GGAAGAAG	2895
2566	UCUUCUUA A UUAGUGUC	568	GACACTAA GGCTAGCTACAACGA ATGGAAGA	2896

2570	CCAUAUUA G UGUCAUCU	569	AGATGACA GGCTAGCTACAACGA TAATATGG	2897
2572	AUAUUAAGU G UCAUCUUG	570	CAAGATGA GGCTAGCTACAACGA ACTAATAT	2898
2575	UUAGUGUC A UCUUGCCU	571	AGGCAAGA GGCTAGCTACAACGA GACACTAA	2899
2580	GUCAUCUU G CCUCCCUA	572	TAGGGAGG GGCTAGCTACAACGA AAGATGAC	2900
2588	GCCUCCCU A CCUCCAC	573	GTGGAAGG GGCTAGCTACAACGA AGGGAGGC	2901
2595	UACCUUCC A CAUGCCCC	574	GGGGCATG GGCTAGCTACAACGA GGAAGGTA	2902
2597	CCUCCAC A UGCCCCAU	575	ATGGGGCA GGCTAGCTACAACGA GTGGAAGG	2903
2599	UUCACAU G CCCCAUGA	576	TCATGGGG GGCTAGCTACAACGA ATGTGGAA	2904
2604	CAUGCCCC A UGACUUGA	577	TCAAGTCA GGCTAGCTACAACGA GGGGCATG	2905
2607	GCCCCAUG A CUUGAUGC	578	GCATCAAG GGCTAGCTACAACGA CATGGGGC	2906
2612	AUGACUUG A UGCAGUUU	579	AAACTGCA GGCTAGCTACAACGA CAAGTCAT	2907
2614	GACUUGAU G CAGUUUUA	580	TAAAACTG GGCTAGCTACAACGA ATCAAGTC	2908
2617	UGAUGCA G UUUUAAUA	581	TATTAAAA GGCTAGCTACAACGA TGCATCAA	2909
2623	CAGUUUUA A UACUUGUA	582	TACAAGTA GGCTAGCTACAACGA TAAAACTG	2910
2625	GUUUUAAU A CUUGUAAU	583	ATTACAAG GGCTAGCTACAACGA ATTAAAAC	2911
2629	UAAUACUU G UAAUUCCC	584	GGGAATTA GGCTAGCTACAACGA AAGTATTA	2912
2632	UACUUGUA A UUCCCCUA	585	TAGGGGAA GGCTAGCTACAACGA TACAAGTA	2913
2641	UUCCCCUA A CCAUAAGA	586	TCTTATGG GGCTAGCTACAACGA TAGGGGAA	2914
2644	CCCUAACC A UAAGAUUU	587	AAATCTTA GGCTAGCTACAACGA GGTTAGGG	2915
2649	ACCAUAAG A UUUACUGC	588	GCAGTAAA GGCTAGCTACAACGA CTTATGGT	2916
2653	UAAGAUUU A CUGCUGCU	589	AGCAGCAG GGCTAGCTACAACGA AAATCTTA	2917
2656	GAUUUACU G CUGCUGUG	590	CACAGCAG GGCTAGCTACAACGA AGTAAATC	2918
2659	UUACUGCU G CUGUGGAU	591	ATCCACAG GGCTAGCTACAACGA AGCAGTAA	2919
2662	CUGCUGCU G UGGAUAUC	592	GATATCCA GGCTAGCTACAACGA AGCAGCAG	2920
2666	UGCUGUGG A UAUCUCCA	593	TGGAGATA GGCTAGCTACAACGA CCACAGCA	2921
2668	CUGUGGAU A UCUCCAUG	594	CATGGAGA GGCTAGCTACAACGA ATCCACAG	2922
2674	AUAUCUCC A UGAAGUUU	595	AAACTTCA GGCTAGCTACAACGA GGAGATAT	2923
2679	UCCAUGAA G UUUUCCCA	596	TGGGAAAA GGCTAGCTACAACGA TTCATGGA	2924
2687	GUUUUCCC A CUGAGUCA	597	TGACTCAG GGCTAGCTACAACGA GGGAAAAC	2925
2692	CCCACUGA G UCA'CAUCA	598	TGATGTGA GGCTAGCTACAACGA TCAGTGGG	2926
2695	ACUGAGUC A CAUCAGAA	599	TTCTGATG GGCTAGCTACAACGA GACTCAGT	2927
2697	UGAGUCAC A UCAGAAU	600	ATTTCTGA GGCTAGCTACAACGA GTGACTCA	2928
2704	CAUCAGAA A UGCCCUC	601	GTAGGGCA GGCTAGCTACAACGA TTCTGATG	2929
2706	UCAGAAAU G CCCUACAU	602	ATGTAGGG GGCTAGCTACAACGA ATTTCTGA	2930
2711	AAUGCCCU A CAUCUUAU	603	ATAAGATG GGCTAGCTACAACGA AGGGCATT	2931
2713	UGCCCUC A UCUUAUUU	604	AAATAAGA GGCTAGCTACAACGA GTAGGGCA	2932
2718	UACAUCUU A UUUUCCUC	605	GAGGAAAA GGCTAGCTACAACGA AAGATGTA	2933
2730	UCCUCAGG G CUCAAGAG	606	CTCTTGAG GGCTAGCTACAACGA CCTGAGGA	2934
2740	UCAAGAGA A UCUGACAG	607	CTGTCAGA GGCTAGCTACAACGA TCTCTTGA	2935
2745	AGAAUCUG A CAGAUACC	608	GGTATCTG GGCTAGCTACAACGA CAGATTCT	2936
2749	UCUGACAG A UACCAUAA	609	TTATGGTA GGCTAGCTACAACGA CTGTCAGA	2937
2751	UGACAGAU A CCAUAAAG	610	CTTTATGG GGCTAGCTACAACGA ATCTGTCA	2938
2754	CAGAUACC A UAAAGGGA	611	TCCCTTTA GGCTAGCTACAACGA GGTATCTG	2939
2762	AUAAAGGG A UUUGACCU	612	AGGTCAAA GGCTAGCTACAACGA CCCTTTAT	2940
2767	GGGAUUUG A CCUAUUA	613	TGATTAGG GGCTAGCTACAACGA CAAATCCC	2941
2772	UUGACCUA A UCACUAAU	614	ATTAGTGA GGCTAGCTACAACGA TAGGTCAA	2942
2775	ACCUAUUC A CUAAUUUJ	615	AAAATTAG GGCTAGCTACAACGA GATTAGGT	2943
2779	AAUCACUA A UUUUCAGG	616	CCTGAAAA GGCTAGCTACAACGA TAGTGATT	2944
2787	AUUUUCAG G UGGUGGCU	617	AGCCACCA GGCTAGCTACAACGA CTGAAAAT	2945
2790	UUCAGGUG G UGGCUGAU	618	ATCAGCCA GGCTAGCTACAACGA CACCTGAA	2946
2793	AGGUGGUG G CUGAUGCU	619	AGCATCAG GGCTAGCTACAACGA CACCACCT	2947
2797	GGUGGCUG A UGCUUUGA	620	TCAAAGCA GGCTAGCTACAACGA CAGCCACC	2948

2799	UGGCUGAU G CUUUGAAC	621	GTTCAAAG GGCTAGCTACAACGA ATCAGCCA	2949
2806	UGC UUUGA A CAUCUCUU	622	AAGAGATG GGCTAGCTACAACGA TCAAAGCA	2950
2808	CUUUGAAC A UCUCUUUG	623	CAAAGAGA GGCTAGCTACAACGA GTTCAAAG	2951
2816	AUCUCUUU G CUGCCCAA	624	TTGGGCAG GGCTAGCTACAACGA AAAGAGAT	2952
2819	UCUUUGCU G CCCAAUCC	625	GGATTGGG GGCTAGCTACAACGA AGCAAAGA	2953
2824	GCUGCCCA A UCCAUIAG	626	CTAATGGA GGCTAGCTACAACGA TGGGCAGC	2954
2828	CCCAAUCC A UUAGCGAC	627	GTCGCTAA GGCTAGCTACAACGA GGATTGGG	2955
2832	AUCCAUIA G CGACAGUA	628	TACTGTCTG GGCTAGCTACAACGA TAATGGAT	2956
2835	CAUUAAGC A CAGUAGGA	629	TCCTACTG GGCTAGCTACAACGA CGCTAATG	2957
2838	UAGCGACA G UAGGAUUU	630	AAATCCTA GGCTAGCTACAACGA TGTCGCTA	2958
2843	ACAGUAGG A UUUUUCAA	631	TTGAAAAA GGCTAGCTACAACGA CCTACTGT	2959
2851	AUUUUUCA A CCCUGGUA	632	TACCAGGG GGCTAGCTACAACGA TGAAAAAT	2960
2857	CAACCCUG G UAUGAAUA	633	TATTCATA GGCTAGCTACAACGA CAGGGTTG	2961
2859	ACCCUGGU A UGAAUAGA	634	TCTATTCA GGCTAGCTACAACGA ACCAGGGT	2962
2863	UGGUAUGA A UAGACAGA	635	TCTGTCTA GGCTAGCTACAACGA TCATACCA	2963
2867	AUGAAUAG A CAGAACCC	636	GGGTCTG GGCTAGCTACAACGA CTATTCAT	2964
2872	UAGACAGA A CCCUAUCC	637	GGATAGGG GGCTAGCTACAACGA TCTGTCTA	2965
2877	AGAACCCU A UCCAGUGG	638	CCACTGGA GGCTAGCTACAACGA AGGGTTCT	2966
2882	CCUAUCCA G UGGAAGGA	639	TCCTFCCA GGCTAGCTACAACGA TGGATAGG	2967
2893	GAAGGAGA A UUUAAUAA	640	TTATTAAA GGCTAGCTACAACGA TCTCCTTC	2968
2898	AGAAUUUA A UAAAGAU	641	TATCTTTA GGCTAGCTACAACGA TAAATTCT	2969
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2907	UAAAGAU A UGCAGAAA	643	TTTCTGCA GGCTAGCTACAACGA TATCTTTA	2971
2909	AAGAUAGU G CAGAAAGA	644	TCTTTCTG GGCTAGCTACAACGA ACTATCTT	2972
2918	CAGAAAGA A UUCCUAG	645	CTAAGGAA GGCTAGCTACAACGA TCTTTCTG	2973
2927	UUCCUAG G UAAUCUUA	646	ATAGATTA GGCTAGCTACAACGA CTAAGGAA	2974
2930	CUUAGGUA A UCUAUAC	647	GTTATAGA GGCTAGCTACAACGA TACCTAAG	2975
2934	GGUAAUCU A UAACUAGG	648	CCTAGTTA GGCTAGCTACAACGA AGATTACC	2976
2937	AAUCUAUA A CUAGGACU	649	AGTCCTAG GGCTAGCTACAACGA TATAGATT	2977
2943	UAACUAGG A CUACUCCU	650	AGGAGTAG GGCTAGCTACAACGA CCTAGTTA	2978
2946	CUAGGACU A CUCCUGGU	651	ACCAGGAG GGCTAGCTACAACGA AGTCCTAG	2979
2953	UACUCCUG G UAACAGUA	652	TACTGTTA GGCTAGCTACAACGA CAGGAGTA	2980
2956	UCCUGGUA A CAGUAAUA	653	TATTACTG GGCTAGCTACAACGA TACCAGGA	2981
2959	UGGUAACA G UAAUACAU	654	ATGTATTA GGCTAGCTACAACGA TGTTACCA	2982
2962	UAACAGUA A UACAUUCC	655	GGAATGTA GGCTAGCTACAACGA TACTGTTA	2983
2964	ACAGUAAU A CAUUCUUA	656	ATGGAATG GGCTAGCTACAACGA ATTACTGT	2984
2966	AGUAAUAC A UUCAUUG	657	CAATGGAA GGCTAGCTACAACGA GTATTACT	2985
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2974	AUUCUUAU G UUUUAGUA	659	TACTAAAA GGCTAGCTACAACGA AATGGAAT	2987
2980	UUGUUUUA G UAACCAGA	660	TCTGGTTA GGCTAGCTACAACGA TAAAACAA	2988
2983	UUUUAGUA A CCAGAAAU	661	ATTTCTGG GGCTAGCTACAACGA TACTAAAA	2989
2990	AACCAGAA A UCUCUUG	662	CATGAAGA GGCTAGCTACAACGA TTCTGGTT	2990
2996	AAAUCUUC A UGCAUUGA	663	TCATTGCA GGCTAGCTACAACGA GAAGATT	2991
2998	AUCUUAU G CAAUGAAA	664	TTTCATTG GGCTAGCTACAACGA ATGAAGAT	2992
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3008	AAUGAAAA A UACUUUAA	666	TTAAAGTA GGCTAGCTACAACGA TTTTCATT	2994
3010	UGAAAAAU A CUUUAAU	667	AATTAAAG GGCTAGCTACAACGA ATTTTTC A	2995
3016	AUACUUUA A UUCAUGAA	668	TTCATGAA GGCTAGCTACAACGA TAAAGTAT	2996
3020	UUUAAUUC A UGAAGCUU	669	AAGCTTCA GGCTAGCTACAACGA GAATTAAA	2997
3025	UUCAUGAA G CUUACUUU	670	AAAGTAAG GGCTAGCTACAACGA TTCATGAA	2998
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3057	AGAGUCUC G CUCUUGUC	675	GACAAGAG GGCTAGCTACAACGA GAGACTCT	3003
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3124	CUUCCCAG G UUCAAGCG	690	CGCTTGAA GGCTAGCTACAACGA CTGGGAAG	3018
3130	AGGUUCAA G CGAUUCUC	691	GAGAATCG GGCTAGCTACAACGA TTGAACCT	3019
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3159	CCUGAGUA G CUGGGAUU	697	AATCCCAG GGCTAGCTACAACGA TACTCAGG	3025
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3174	UUACAGGC G UGUGCACU	701	AGTGCACA GGCTAGCTACAACGA GCCTGTAA	3029
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3178	AGGCGUGU G CACUACAC	703	GTGTAGTG GGCTAGCTACAACGA ACACGCCT	3031
3180	GCGUGUGC A CUACACUC	704	GAGTGTAG GGCTAGCTACAACGA GCACACGC	3032
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3233	ACUGUUG G CCAGGCUG	715	CAGCCTGG GGCTAGCTACAACGA CAACAGGT	3043
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3242	CCAGGCUG G UCUCGAAC	717	GTTTCGAGA GGCTAGCTACAACGA CAGCCTGG	3045
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3256	AACUCCUG A CCUCAAGU	719	ACTTGAGG GGCTAGCTACAACGA CAGGAGTT	3047
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3361	CGCACAAG G CACUGGGU	745	ACCCAGTG GGCTAGCTACAACGA CTGTGCGG	3073
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3439	AAUAUCUU A CUAAGGCC	766	GGCCTTAG GGCTAGCTACAACGA AAGATATT	3094
3445	UUACUAAG G CCUUUGGU	767	ACCAAAGG GGCTAGCTACAACGA CTTAGTAA	3095
3452	GGCCUUUG G UAUACGAC	768	GTCGTATA GGCTAGCTACAACGA CAAAGGCC	3096
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3456	UUUGGUUA A CGACCCAG	770	CTGGGTCG GGCTAGCTACAACGA ATACCAA	3098
3459	GGUAUACG A CCCAGAGA	771	TCTCTGGG GGCTAGCTACAACGA CGTATACC	3099
3467	ACCCAGAG A UAACACGA	772	TCGTGTTA GGCTAGCTACAACGA CTCTGGGT	3100
3470	GAGAGUA A CACGAUGC	773	GCATCGTG GGCTAGCTACAACGA TATCTCTG	3101
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3477	AACACGAU G CGUAUUUU	776	AAAATACG GGCTAGCTACAACGA ATCGTGT	3104

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3528	GCUCUAUA A UUGUUUUG	787	CAAAACAA GGCTAGCTACAACGA TATAGAGC	3115
3531	CUAUAUUU G UUUUGCUA	788	TAGCAAAA GGCTAGCTACAACGA AATTATAG	3116
3536	AUUGUUUU G CUACGAUU	789	AATCGTAG GGCTAGCTACAACGA AAAACAAT	3117
3539	GUUUUGCU A CGAUUCCA	790	TGGAATCG GGCTAGCTACAACGA AGCAAAAC	3118
3542	UUGCUACG A UUCCACUG	791	CAGTGGAA GGCTAGCTACAACGA CGTAGCAA	3119
3547	ACGAUUC A CUGAAACU	792	AGTTTCAG GGCTAGCTACAACGA GGAATCGT	3120
3553	CCACUGAA A CUCUUCGA	793	TCGAAGAG GGCTAGCTACAACGA TTCAGTGG	3121
3561	ACUCUUCG A UCAAGCUA	794	TAGCTTGA GGCTAGCTACAACGA CGAAGAGT	3122
3566	UCGAUCAA G CUACUUUA	795	TAAAGTAG GGCTAGCTACAACGA TTGATCGA	3123
3569	AUCAAGCU A CUUUAUGU	796	ACATAAAG GGCTAGCTACAACGA AGCTTGAT	3124
3574	GCUACUUU A UGUAAAUC	797	GATTTACA GGCTAGCTACAACGA AAAGTAGC	3125
3576	UACUUUAU G UAAAUAC	798	GTGATTTA GGCTAGCTACAACGA ATAAAGTA	3126
3580	UUUAUGUA A UCACUUA	799	TGAAGTGA GGCTAGCTACAACGA TTACATAA	3127
3583	UGUAAAUC A CUUCAUUG	800	CAATGAAG GGCTAGCTACAACGA GATTTACA	3128
3588	AUCACUUC A UUGUUUUA	801	TAAAACAA GGCTAGCTACAACGA GAAGTGAT	3129
3591	ACUUCAUU G UUUUAAAG	802	CTTTAAAA GGCTAGCTACAACGA AATGAAGT	3130
3602	UUAAAGGA A UAAACUUG	803	CAAGTTTA GGCTAGCTACAACGA TCCTTTAA	3131
3606	AGGAUUA A CUUGAUUA	804	TAATCAAG GGCTAGCTACAACGA TTATTCCT	3132
3611	UAAACUUG A UUAUUAUG	805	CAATATAA GGCTAGCTACAACGA CAAGTTTA	3133
3614	ACUUGAUU A UAUUGUUU	806	AAACAATA GGCTAGCTACAACGA AATCAAGT	3134
3616	UUGAUUAU A UUGUUUUU	807	AAAAACAA GGCTAGCTACAACGA ATAATCAA	3135
3619	AUUAUAUU G UUUUUUUA	808	TAAAAAAA GGCTAGCTACAACGA AATATAAT	3136
3627	GUUUUUUU A UUUGGCAU	809	ATGCCAAA GGCTAGCTACAACGA AAAAAAAC	3137
3632	UUUAUUUG G CAUAACUG	810	CAGTTATG GGCTAGCTACAACGA CAAATAAA	3138
3634	UAUUUGGC A UAACUGUG	811	CACAGTTA GGCTAGCTACAACGA GCCAAATA	3139
3637	UUGGCAUA A CUGUGAUU	812	AATCACAG GGCTAGCTACAACGA TATGCCAA	3140
3640	GCAUAACU G UGAUUCUU	813	AAGAATCA GGCTAGCTACAACGA AGTTATGC	3141
3643	UAACUGUG A UUCUUUUA	814	TAAAAGAA GGCTAGCTACAACGA CACAGTTA	3142
3654	CUUUUAGG A CAAUUAU	815	AGTAATTG GGCTAGCTACAACGA CCTAAAAG	3143
3657	UUAGGACA A UUACUGUA	816	TACAGTAA GGCTAGCTACAACGA TGTCTTAA	3144
3660	GGACAAUU A CUGUACAC	817	GTGTACAG GGCTAGCTACAACGA AATFGTCC	3145
3663	CAUUUAU G UACACAUU	818	AATGTGTA GGCTAGCTACAACGA AGTAATTG	3146
3665	AUUACUGU A CACAUUAA	819	TTAATGTG GGCTAGCTACAACGA ACAGTAAT	3147
3667	UACUGUAC A CAUUAAGG	820	CCTTAATG GGCTAGCTACAACGA GTACAGTA	3148
3669	CUGUACAC A UUAAGGUG	821	CACCTTAA GGCTAGCTACAACGA GTGTACAG	3149
3675	ACAUUAAG G UGUUGUC	822	GACATACA GGCTAGCTACAACGA CTTAATGT	3150
3677	AUUAAGGU G UAUUGUCAG	823	CTGACATA GGCTAGCTACAACGA ACCTTAAT	3151
3679	UAAGGUGU A UGUCAGAU	824	ATCTGACA GGCTAGCTACAACGA ACACCTTA	3152
3681	AGGUGUAU G UCAGUAU	825	ATATCTGA GGCTAGCTACAACGA ATACACCT	3153
3686	UAUGUCAG A UAUUCAUA	826	TATGAATA GGCTAGCTACAACGA CTGACATA	3154
3688	UGUCAGAU A UUCAUAUU	827	AATATGAA GGCTAGCTACAACGA ATCTGACA	3155
3692	AGAUUAUC A UAUUGACC	828	GGTCAATA GGCTAGCTACAACGA GAATATCT	3156

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3698	UCAUAUUG A CCCAAUUG	830	CATTTGGG GGCTAGCTACAACGA CAATATGA	3158
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3706	ACCCAAAU G UGUAAUUAU	832	ATATTACA GGCTAGCTACAACGA ATTTGGGT	3160
3708	CCAAUUGU G UAAUAUUC	833	GAATATTA GGCTAGCTACAACGA ACATTTGG	3161
3711	AAUGUGUA A UAUUCCAG	834	CTGGAATA GGCTAGCTACAACGA TACACATT	3162
3713	UGUGUAAU A UUCCAGUU	835	AACTGGAA GGCTAGCTACAACGA ATTACACA	3163
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3728	UUUUCUCU G CAUAAGUA	837	TACTTATG GGCTAGCTACAACGA AGAGAAAA	3165
3730	UUCUCUGC A UAAGUAAU	838	ATTACTTA GGCTAGCTACAACGA GCAGAGAA	3166
3734	CUGCAUAA G UAAUAAAA	839	TTTAATTA GGCTAGCTACAACGA TTATGCAG	3167
3737	CAUAAGUA A UUAAAAUA	840	TATTTTAA GGCTAGCTACAACGA TACTTATG	3168
3743	UAAUAAAA A UAUACUUA	841	TAAGTATA GGCTAGCTACAACGA TTTAATTA	3169
3745	AUUAAAAU A UACUAAAA	842	TTTAAGTA GGCTAGCTACAACGA ATTTTAAT	3170
3747	UAAAAUAU A CUUAAAAA	843	TTTTTAAG GGCTAGCTACAACGA ATATTTTA	3171
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3759	AAAAAUUA A UAGUUUUA	845	TAAACTTA GGCTAGCTACAACGA TAATTTTT	3173
3762	AAUUAUAU G UUUUAUCU	846	AGATAAAA GGCTAGCTACAACGA TATTAATT	3174
3767	AUAGUUUU A UCUGGGUA	847	TACCCAGA GGCTAGCTACAACGA AAAACTAT	3175
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3775	AUCUGGGU A CAAUAAAA	849	TTTATTTG GGCTAGCTACAACGA ACCCAGAT	3177
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3786	AAUAAACA G UGCCUGAA	852	TTCAGGCA GGCTAGCTACAACGA TGTTTATT	3180
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3806	GUUCACAG A CAAGGGAA	857	TTCCCTTG GGCTAGCTACAACGA CTGTGAAC	3185
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3832	UAAAAAUC A CUAUGAUU	862	AATCATAG GGCTAGCTACAACGA GATTTTTA	3190
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3838	UCACUAUG A UUUCUGAA	864	TTCAGAAA GGCTAGCTACAACGA CATAGTGA	3192
3846	AUUUCUGA A UUGCUAUG	865	CATAGCAA GGCTAGCTACAACGA TCAGAAAT	3193
3849	UCUGAAUU G CUAUGUGA	866	TCACATAG GGCTAGCTACAACGA AATTCAGA	3194
3852	GAAUUGCU A UGUGAAAC	867	GTTTCACA GGCTAGCTACAACGA AGCAATTC	3195
3854	AUUGCUAU G UGAAACUA	868	TAGTTTCA GGCTAGCTACAACGA ATAGCAAT	3196
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3862	GUGAAACU A CAGAUUUU	870	AAGATCTG GGCTAGCTACAACGA AGTTTCAC	3198
3866	AACUACAG A UCUUUGGA	871	TCCAAAGA GGCTAGCTACAACGA CTGTAGTT	3199
3875	UCUUUGGA A CACUGUUU	872	AAACAGTG GGCTAGCTACAACGA TCCAAAGA	3200
3877	UUUGGAAC A CUGUUUAG	873	CTAAACAG GGCTAGCTACAACGA GTTCCAAA	3201
3880	GGAACACU G UUUAGGUA	874	TACCTAAA GGCTAGCTACAACGA AGTGTTCC	3202
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3891	UAGGUAGG G UGUUAGA	876	TCCTTAACA GGCTAGCTACAACGA CCTACCTA	3204
3893	GGUAGGGU G UUAAGACU	877	AGTCTTAA GGCTAGCTACAACGA ACCCTACC	3205
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3906	GACUUGAC A CAGUACCU	880	AGGTACTG GGCTAGCTACAACGA GTCAAGTC	3208

3909	UUGACACA G UACCUCGU	881	ACGAGGTA GGCTAGCTACAACGA TGTGTCAA	3209
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3916	AGUACCUC G UUUUACA	883	TGTAGAAA GGCTAGCTACAACGA GAGGTACT	3211
3922	UCGUUUCU A CACAGAGA	884	TCTCTGTG GGCTAGCTACAACGA AGAAACGA	3212
3924	GUUUCUAC A CAGAGAAA	885	TTTCTCTG GGCTAGCTACAACGA GTAGAAAC	3213
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3939	AAGAAUG G CCAUACU	887	AAGTATGG GGCTAGCTACAACGA CATTTTCT	3215
3942	AAAUGGCC A UACUUCAG	888	CTGAAGTA GGCTAGCTACAACGA GGCCATTT	3216
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3961	ACUGCAGU G CUUAUGAG	893	CTCATAAG GGCTAGCTACAACGA ACTGCAGT	3221
3965	CAGUGCUU A UGAGGGGA	894	TCCCCTCA GGCTAGCTACAACGA AAGCACTG	3222
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3981	AUAUUUAG G CCUCUUGA	897	TCAAGAGG GGCTAGCTACAACGA CTAAATAT	3225
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4125	ACCUAAUU A UUACAGCC	925	GGCTGTAA GGCTAGCTACAACGA AATTAGGT	3253
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4537	AGAUUCC A UUUGUCA	1018	TTGACAAA GGCTAGCTACAACGA GGAAATCT	3346
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4552	AAAAAGUA A UGAUUUCU	1021	AGAAATCA GGCTAGCTACAACGA TACTTTTT	3349
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4566	UCUUGAUA A UUGUGUAG	1024	CTACACAA GGCTAGCTACAACGA TATCAAGA	3352
4569	UGAUAAUU G UGUAGUGA	1025	TCACTACA GGCTAGCTACAACGA AATTATCA	3353
4571	AUAAUUGU G UAGUGAAU	1026	ATTCACCTA GGCTAGCTACAACGA ACAATTAT	3354
4574	AUUGUGUA G UGAAUGUU	1027	AACATTCA GGCTAGCTACAACGA TACACAAT	3355
4578	UGUAGUGA A UGUUUUU	1028	AAAAACA GGCTAGCTACAACGA TCACTACA	3356
4580	UAGUGAAU G UUUUUUAG	1029	CTAAAAAA GGCTAGCTACAACGA ATTCACCTA	3357
4590	UUUUUAGA A CCCAGCAG	1030	CTGCTGGG GGCTAGCTACAACGA TCTAAAAA	3358
4595	AGAACCCA G CAGUUACC	1031	GGTAACTG GGCTAGCTACAACGA TGGGTTCT	3359
4598	ACCCAGCA G UUACCUUG	1032	CAAGGTAA GGCTAGCTACAACGA TGCTGGGT	3360
4601	CAGCAGUU A CCUUGAAA	1033	TTTCAAGG GGCTAGCTACAACGA AACTGCTG	3361
4610	CCUUGAAA G CUGAAUUU	1034	AAATTCAAG GGCTAGCTACAACGA TTTCAAGG	3362
4615	AAAGCUGA A UUAUAUU	1035	AATATAAA GGCTAGCTACAACGA TCAGCTTT	3363
4619	CUGAAUUU A UAUUUAGU	1036	ACTAAATA GGCTAGCTACAACGA AAATTCAAG	3364

4621	GAAUUUAU A UUUAGUAA	1037	TTACTAAA GGCTAGCTACAACGA ATAAATTC	3365
4626	UAUAUUUA G UAACUUCU	1038	AGAAGTTA GGCTAGCTACAACGA TAAATATA	3366
4629	AUUUAGUA A CUUCUGUG	1039	CACAGAAG GGCTAGCTACAACGA TACTAAAT	3367
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4637	ACUUCUGU G UUAUACU	1041	AGTATTAA GGCTAGCTACAACGA ACAGAAGT	3369
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4643	GUGUUAAU A CUGGAUAG	1043	CTATCCAG GGCTAGCTACAACGA ATTAACAC	3371
4648	AAUACUGG A UAGCAUGA	1044	TCATGCTA GGCTAGCTACAACGA CCAGTATT	3372
4651	ACUGGAUA G CAUGAAUU	1045	AATTCATG GGCTAGCTACAACGA TATCCAGT	3373
4653	UGGAUAGC A UGAAUUCU	1046	AGAATTCA GGCTAGCTACAACGA GCTATCCA	3374
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4776	GUUUAAUA G UUUGAAGU	1072	ACTTCAA GGCTAGCTACAACGA TATTAAAC	3400
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5435	UGUUAGAA G UAUCCUUU	1237	AAAGGATA GGCTAGCTACAACGA TTCTAACA	3565
5437	UUAGAAGU A UCCUUUUA	1238	TAAAAGGA GGCTAGCTACAACGA ACTTCTAA	3566
5445	AUCCUUUU A UUUUCUAA	1239	TTAGAAAA GGCTAGCTACAACGA AAAAGGAT	3567
5457	UCUAAAAG G UGCUGUGG	1240	CCACAGCA GGCTAGCTACAACGA CTTTTAGA	3568
5459	UAAAAGGU G CUGUGGAU	1241	ATCCACAG GGCTAGCTACAACGA ACCTTTTA	3569
5462	AAGGUGCU G UGGAUUAU	1242	AATATCCA GGCTAGCTACAACGA AGCACCTT	3570
5466	UGCUGUGG A UAUUAUGU	1243	ACATAATA GGCTAGCTACAACGA CCACAGCA	3571
5468	CUGUGGAU A UUAUGUAA	1244	TTACATAA GGCTAGCTACAACGA ATCCACAG	3572

5471	UGGAUUAU A UGUAAAGG	1245	CCTTTACA GGCTAGCTACAACGA AATATCCA	3573
5473	GAUUAUUAU G UAAAGGCG	1246	CGCCTTTA GGCTAGCTACAACGA ATAATATC	3574
5479	AUGUAAAG G CGUGUUUG	1247	CAAACACG GGCTAGCTACAACGA CTTTACAT	3575
5481	GUAAAGGC G UGUUUGCU	1248	AGCAAACA GGCTAGCTACAACGA GCCTTTAC	3576
5483	AAAGGCGU G UUUGCUUA	1249	TAAGCAAA GGCTAGCTACAACGA ACGCCTTT	3577
5487	GCGUGUUU G CUUAAACA	1250	TGTTTAAG GGCTAGCTACAACGA AAACACGC	3578
5493	UUGCUUAA A CAAUUUUC	1251	GAAAAATG GGCTAGCTACAACGA TTAAGCAA	3579
5496	CUUAAACA A UUUUCCAU	1252	ATGGAAAA GGCTAGCTACAACGA TGTTTAAG	3580
5503	AAUUUUC A UAUUUAGA	1253	TCTAAATA GGCTAGCTACAACGA GGAAAAAT	3581
5505	UUUCCAU A UUUAGAAG	1254	CTTCTAAA GGCTAGCTACAACGA ATGGAAAA	3582
5513	AUUUAGAA G UAGAUGCA	1255	TGCATCTA GGCTAGCTACAACGA TTCTAAAT	3583
5517	AGAAGUAG A UGCAAAAC	1256	GTTTGTGA GGCTAGCTACAACGA CTACTTCT	3584
5519	AAGUAGAU G CAAAACAA	1257	TTGTTTGG GGCTAGCTACAACGA ATCTACTT	3585
5524	GAUGCAAA A CAAUCUG	1258	CAGATTTG GGCTAGCTACAACGA TTTGCATC	3586
5528	CAAAACAA A UCUGCCUU	1259	AAGGCAGA GGCTAGCTACAACGA TTGTTTGG	3587
5532	ACAAUCU G CCUUUAUG	1260	CATAAAGG GGCTAGCTACAACGA AGATTTGT	3588
5538	CUGCCUUU A UGACAAA	1261	TTTGTGCA GGCTAGCTACAACGA AAAGGCAG	3589
5541	CCUUUAUG A CAAAAAA	1262	TTTTTTGG GGCTAGCTACAACGA CATAAAGG	3590
5549	ACAAAAA A UAGGAUAA	1263	TTATCCTA GGCTAGCTACAACGA TTTTTTGT	3591
5554	AAAUAAGG A UAACAUUA	1264	TAATGTTA GGCTAGCTACAACGA CCTATTTT	3592
5557	AUAGGAUA A CAUUAUUU	1265	AAATAATG GGCTAGCTACAACGA TATCCTAT	3593
5559	AGGAUAAC A UUAUUUAU	1266	ATAAATAA GGCTAGCTACAACGA GTTATCCT	3594
5562	AUAACAUU A UUAUUUA	1267	TAAATAAA GGCTAGCTACAACGA AATGTTAT	3595
5566	CAUUAUUU A UUAUUUC	1268	GAAATAAA GGCTAGCTACAACGA AAATAATG	3596
5570	AUUUAUUU A UUUCUUU	1269	AAAGGAAA GGCTAGCTACAACGA AAATAAAT	3597
5580	UUCCUUUU A UCAUAAG	1270	CTTATTGA GGCTAGCTACAACGA AAAAGGAA	3598
5584	UUUAUCA A UAAGGUA	1271	TTACCTTA GGCTAGCTACAACGA TGATAAAA	3599
5589	UCAUAAG G UAAUUGAU	1272	ATCAATTA GGCTAGCTACAACGA CTTATTGA	3600
5592	AUAAGGUA A UUGAUACA	1273	TGTATCAA GGCTAGCTACAACGA TACCTTAT	3601
5596	GGUAAUUG A UACACAAC	1274	GTTGTGTA GGCTAGCTACAACGA CAATTACC	3602
5598	UAAUUGAU A CACAACAG	1275	CTGTTGTG GGCTAGCTACAACGA ATCAATTA	3603
5600	AUUGAUAC A CAACAGGU	1276	ACCTGTTG GGCTAGCTACAACGA GTATCAAT	3604
5603	GAUACACA A CAGGUGAC	1277	GTCACCTG GGCTAGCTACAACGA TGTGTATC	3605
5607	CACAACAG G UGACUUGG	1278	CCAAGTCA GGCTAGCTACAACGA CTGTTGTG	3606
5610	AACAGGUG A CUUGGUUU	1279	AAACCAAG GGCTAGCTACAACGA CACCTGTT	3607
5615	GUGACUUG G UUUUAGGC	1280	GCCTAAAA GGCTAGCTACAACGA CAAGTCAC	3608
5622	GGUUUUAG G CCCAAAGG	1281	CCTTTGGG GGCTAGCTACAACGA CTAAAAACC	3609
5630	GCCCAAAG G UAGCAGCA	1282	TGCTGCTA GGCTAGCTACAACGA CTTTGGGC	3610
5633	CAAAGGUA G CAGCAGCA	1283	TGCTGCTG GGCTAGCTACAACGA TACCTTTG	3611
5636	AGGUAGCA G CAGCAACA	1284	TGTTGCTG GGCTAGCTACAACGA TGCTACCT	3612
5639	UAGCAGCA G CAACAUUA	1285	TAATGTTG GGCTAGCTACAACGA TGCTGCTA	3613
5642	CAGCAGCA A CAUUAUA	1286	TATTAATG GGCTAGCTACAACGA TGCTGCTG	3614
5644	GCAGCAAC A UUAUAUA	1287	ATTATTAA GGCTAGCTACAACGA GTTGCTGC	3615
5648	CAACAUUA A UAAUGGAA	1288	TTCCATTA GGCTAGCTACAACGA TAATGTTG	3616
5651	CAUUAUA A UGGAAUA	1289	TATTTCCA GGCTAGCTACAACGA TATTAATG	3617
5657	UAAUGGAA A UAAUUGAA	1290	TTCAATTA GGCTAGCTACAACGA TTCCATTA	3618
5660	UGGAAUA A UUGAAUAG	1291	CTATTCAA GGCTAGCTACAACGA TATTTCCA	3619
5665	AUAUUGA A UAGUUAGU	1292	ACTAACTA GGCTAGCTACAACGA TCAATTAT	3620
5668	AUUGAAUA G UUAGUUUA	1293	ATAACTAA GGCTAGCTACAACGA TATTCAAT	3621
5672	AAUAGUUA G UUAUGUAU	1294	ATACATAA GGCTAGCTACAACGA TAACTATT	3622
5675	AGUAGUU A UGUAGUU	1295	AACATACA GGCTAGCTACAACGA AACTAACT	3623
5677	UUAGUUUA G UAUGUUA	1296	TTAACATA GGCTAGCTACAACGA ATAATAA	3624

5679	AGUUAUGU A UGUUAAUG	1297	CATTAACA GGCTAGCTACAACGA ACATAACT	3625
5681	UUAUGUUAU G UUAUGCC	1298	GGCATTAA GGCTAGCTACAACGA ATACATAA	3626
5685	GUAUGUUA A UGCCAGUC	1299	GACTGGCA GGCTAGCTACAACGA TAACATAC	3627
5687	AUGUUAU G CCAGUCAC	1300	GTGACTGG GGCTAGCTACAACGA ATTAACAT	3628
5691	UUAUGCCA G UCACCAGC	1301	GCTGGTGA GGCTAGCTACAACGA TGGCATT	3629
5694	UGCCAGUC A CCAGCAGG	1302	CCTGCTGG GGCTAGCTACAACGA GACTGGCA	3630
5698	AGUCACCA G CAGGCUAU	1303	ATAGCCTG GGCTAGCTACAACGA TGGTGACT	3631
5702	ACCAGCAG G CUAUUUCA	1304	TGAAATAG GGCTAGCTACAACGA CTGCTGGT	3632
5705	AGCAGGCU A UUCAAGG	1305	CCTGAAA GGCTAGCTACAACGA AGCCTGCT	3633
5713	AUUUCAAG G UCAGAAGU	1306	ACTTCTGA GGCTAGCTACAACGA CTTGAAAT	3634
5720	GGUCAGAA G UAAUGACU	1307	AGTCATTA GGCTAGCTACAACGA TTCTGACC	3635
5723	CAGAAGUA A UGACUCCA	1308	TGGAGTCA GGCTAGCTACAACGA TACTTCTG	3636
5726	AAGUAAUG A CUCCAUAU	1309	GTATGGAG GGCTAGCTACAACGA CATTACTT	3637
5731	AUGACUCC A UACAUUU	1310	AATATGTA GGCTAGCTACAACGA GGAGTCAT	3638
5733	GACUCCA U CAUAUUUAU	1311	ATAATATG GGCTAGCTACAACGA ATGGAGTC	3639
5735	CUCCAUAU A UAUUAUUU	1312	AAATAATA GGCTAGCTACAACGA GTATGGAG	3640
5737	CCAUACAU A UUAUUUAU	1313	ATAAATAA GGCTAGCTACAACGA ATGTATGG	3641
5740	UACAUUUU A UUUUUUUC	1314	GAAATAAA GGCTAGCTACAACGA AATATGTA	3642
5744	UAUUUUUU A UUUUUAUA	1315	TATAGAAA GGCTAGCTACAACGA AAATAATA	3643
5750	UUUUUUUCU A UAACUACA	1316	TGTAGTTA GGCTAGCTACAACGA AGAAATAA	3644
5753	UUUCUAUA A CUACAUUU	1317	AAATGTAG GGCTAGCTACAACGA TATAGAAA	3645
5756	CUUAACU A CAUUUAAA	1318	TTTAAATG GGCTAGCTACAACGA AGTTATAG	3646
5758	AUAACUAC A UUUAAAUC	1319	GATTTAAA GGCTAGCTACAACGA GTAGTTAT	3647
5764	ACAUUUAA A UCAUUACC	1320	GGTAATGA GGCTAGCTACAACGA TTAAATGT	3648
5767	UUUAAAUC A UUACCAGG	1321	CCTGGTAA GGCTAGCTACAACGA GATTTAAA	3649

Input Sequence = NM_004985. Cut Site = R/Y

Arm Length = 8. Core Sequence = GGCTAGCTACAACGA

NM_004985 (Homo sapiens v-Ki-ras2 Kirsten rat sarcoma 2 viral oncogene homolog (KRas2), mRNA; 5775 nt)

Table III: Human H-Ras DNzyme and Target molecules

Pos	Substrate	Seq ID	DNzyme	Seq ID
9	GGAUCCCA G CCUUUCCC	1322	GGGAAAGG GGCTAGCTACAACGA TGGGATCC	3650
20	UUUCCCCA G CCCGUAGC	1323	GCTACGGG GGCTAGCTACAACGA TGGGGAAA	3651
24	CCCAGCCC G UAGCCCCG	1324	CGGGGCTA GGCTAGCTACAACGA GGGCTGGG	3652
27	AGCCCGUA G CCCCGGGA	1325	TCCCGGGG GGCTAGCTACAACGA TACGGGCT	3653
35	GCCCCGGG A CCUCCGCG	1326	CGCGGAGG GGCTAGCTACAACGA CCCGGGGC	3654
41	GGACCUCC G CGGUGGGC	1327	GCCCACCG GGCTAGCTACAACGA GGAGGTCC	3655
44	CCUCCGCG G UGGGCGGC	1328	GCCGCCCA GGCTAGCTACAACGA CGCGGAGG	3656
48	CGCGGUGG G CGGCGCCG	1329	CGGCGCCG GGCTAGCTACAACGA CCACCGCG	3657
51	GGUGGGCG G CGCCGCGC	1330	GCGCGCGG GGCTAGCTACAACGA CGCCACC	3658
53	UGGGCGGC G CCGCGCUG	1331	CAGCGCGG GGCTAGCTACAACGA GCCGCCCA	3659
56	GCGGCGCC G CGCUGCCG	1332	CGGCAGCG GGCTAGCTACAACGA GCGCGCCG	3660
58	GGCGCCGC G CUGCCGGC	1333	GCCGGCAG GGCTAGCTACAACGA GCGGCGCC	3661
61	GCCGCGCU G CCGGCGCA	1334	TGCGCCCG GGCTAGCTACAACGA AGCGCGGC	3662
65	CGCUGCCG G CGCAGGGA	1335	TCCCTGCG GGCTAGCTACAACGA CGGCAGCG	3663
67	CUGCCGGC G CAGGGAGG	1336	CCTCCCTG GGCTAGCTACAACGA GCCGGCAG	3664
76	CAGGGAGG G CCUCUGGU	1337	ACCAGAGG GGCTAGCTACAACGA CCTCCCTG	3665
83	GGCCUCUG G UGCACCGG	1338	CCGGTGCA GGCTAGCTACAACGA CAGAGGCC	3666
85	CCUCUGGU G CACCGGCA	1339	TGCCGGTG GGCTAGCTACAACGA ACCAGAGG	3667
87	UCUGGUGC A CCGGCACC	1340	GGTGCCCG GGCTAGCTACAACGA GCACCAGA	3668
91	GUGCACCG G CACCGCUG	1341	CAGCGGTG GGCTAGCTACAACGA CGGTGCAC	3669
93	GCACCGGC A CCGCUGAG	1342	CTCAGCGG GGCTAGCTACAACGA GCCGGTGC	3670
96	CCGGCACC G CUGAGUCG	1343	CGACTCAG GGCTAGCTACAACGA GGTGCCCG	3671
101	ACCGCUGA G UCGGGUUC	1344	GAACCCGA GGCTAGCTACAACGA TCAGCGGT	3672
106	UGAGUCGG G UUCUCUCG	1345	CGAGAGAA GGCTAGCTACAACGA CCGACTCA	3673
114	GUUCUCUC G CCGGCCUG	1346	CAGGCCCG GGCTAGCTACAACGA GAGAGAAC	3674
118	UCUCGCCG G CCUGUUC	1347	GGAACAGG GGCTAGCTACAACGA CGGCGAGA	3675
122	GCCGGCCU G UUCCCGGG	1348	CCCGGGAA GGCTAGCTACAACGA AGGCCGGC	3676
134	CCGGGAGA G CCCGGGGC	1349	GCCCCGGG GGCTAGCTACAACGA TCTCCCGG	3677
141	AGCCCGGG G CCCUGCUC	1350	GAGCAGGG GGCTAGCTACAACGA CCCGGGCT	3678
146	GGGGCCCU G CUCGGAGA	1351	TCTCCGAG GGCTAGCTACAACGA AGGGCCCC	3679
154	GCUCGGAG A UGCCGCC	1352	GGGCGGCA GGCTAGCTACAACGA CTCCGAGC	3680
156	UCGAGAU G CGGCCCG	1353	CGGGGCGG GGCTAGCTACAACGA ATCTCCGA	3681
159	GAGAUGCC G CCCCGGGC	1354	GCCCCGGG GGCTAGCTACAACGA GGCATCTC	3682
166	CGCCCCGG G CCCCCAGA	1355	TCTGGGGG GGCTAGCTACAACGA CCGGGGCG	3683
174	GCCCCCAG A CACCGGCU	1356	AGCCGGTG GGCTAGCTACAACGA CTGGGGGC	3684
176	CCCCAGAC A CCGGCUC	1357	GGAGCCCG GGCTAGCTACAACGA GTCTGGGG	3685
180	AGACACCG G CUCCUGG	1358	CCAGGGAG GGCTAGCTACAACGA CGGTGTCT	3686
188	GCUCCUG G CCUCCUC	1359	GAGGAAGG GGCTAGCTACAACGA CAGGGAGC	3687
199	UUCUCGA G CAACCCCG	1360	CGGGGTTG GGCTAGCTACAACGA TCGAGGAA	3688
202	CUCGAGCA A CCCCAGC	1361	GCTCGGGG GGCTAGCTACAACGA TGCTCGAG	3689
209	AACCCCGA G CUCGGCUC	1362	GAGCCGAG GGCTAGCTACAACGA TCGGGGTT	3690
214	CGAGCUCG G CUCCGGUC	1363	GACCGGAG GGCTAGCTACAACGA CGAGCTCG	3691
220	CGGCUCCG G UCUCAGC	1364	GCTGGAGA GGCTAGCTACAACGA CGGAGCCG	3692
227	GGUCUCCA G CCAAGCCC	1365	GGGCTTGG GGCTAGCTACAACGA TGGAGACC	3693
232	CCAGCCAA G CCCAACCC	1366	GGGTTGGG GGCTAGCTACAACGA TTGGCTGG	3694
237	CAAGCCCA A CCCCAGAG	1367	TCTCGGGG GGCTAGCTACAACGA TGGGCTTG	3695
247	CCCAGAG G CCGCGGCC	1368	GGCCGCGG GGCTAGCTACAACGA CTCTCGGG	3696
250	GAGAGGCC G CGGCCUA	1369	TAGGGCCG GGCTAGCTACAACGA GGCCTCTC	3697

253	AGGCCGCG G CCCUACUG	1370	CAGTAGGG GGCTAGCTACAACGA CGCGGCCT	3698
258	GCGGCCCU A CUGGCUCC	1371	GGAGCCAG GGCTAGCTACAACGA AGGGCCGC	3699
262	CCCUACUG G CUCCGCCU	1372	AGGCGGAG GGCTAGCTACAACGA CAGTAGGG	3700
267	CUGGCUCC G CCUCCGCG	1373	GCGGGAGG GGCTAGCTACAACGA GGAGCCAG	3701
274	CGCCUCCC G CGUUGCUC	1374	GAGCAACG GGCTAGCTACAACGA GGGAGGCG	3702
276	CCUCCGCG G UUGCUC	1375	GGGAGCAA GGCTAGCTACAACGA GCGGGAGG	3703
279	CCCGCGUU G CUCCCGGA	1376	TCCGGGAG GGCTAGCTACAACGA AACGCGGG	3704
289	UCCCGGAA G CCCCGCCC	1377	GGGCGGGG GGCTAGCTACAACGA TTCCGGGA	3705
294	GAAGCCCC G CCCGACCG	1378	CGGTCTGG GGCTAGCTACAACGA GGGGCTTC	3706
299	CCCGCCCG A CCGCGGCU	1379	AGCCGCGG GGCTAGCTACAACGA CGGGCGGG	3707
302	GCCCGACC G CGGCUCCU	1380	AGGAGCCG GGCTAGCTACAACGA GGTCTGGC	3708
305	CGACCGCG G CUCCUGAC	1381	GTCAGGAG GGCTAGCTACAACGA CGCGGTCTG	3709
312	GGCUCCUG A CAGACGGG	1382	CCCGTCTG GGCTAGCTACAACGA CAGGAGCC	3710
316	CCUGACAG A CGGGCGCG	1383	GCGGCCCG GGCTAGCTACAACGA CTGTCAAG	3711
320	ACAGACGG G CCGCUCAG	1384	CTGAGCGG GGCTAGCTACAACGA CCGTCTGT	3712
323	GACGGGCC G CUCAGCCA	1385	TGGCTGAG GGCTAGCTACAACGA GGCCCGTC	3713
328	GCCGCUCA G CCAACCGG	1386	CCGTTGGG GGCTAGCTACAACGA TGAGCGGC	3714
332	CUCAGCCA A CCGGGGUG	1387	CACCCCGG GGCTAGCTACAACGA TGGCTGAG	3715
338	CAACCGGG G UGGGGCGG	1388	CCGCCCA GGCTAGCTACAACGA CCCGGTTG	3716
343	GGGGUGGG G CGGGGCCC	1389	GGGCCCGG GGCTAGCTACAACGA CCCACCCC	3717
348	GGGGCGGG G CCCGAUGG	1390	CCATCGGG GGCTAGCTACAACGA CCCGCCCC	3718
353	GGGGCCCG A UGGCGCGC	1391	GCGCGCCA GGCTAGCTACAACGA CGGGCCCC	3719
356	GCCCGAUG G CGCGCAGC	1392	GCTGCGCG GGCTAGCTACAACGA CATCGGGC	3720
358	CCGAUGGC G CGCAGCCA	1393	TGGCTGCG GGCTAGCTACAACGA GCCATCGG	3721
360	GAUGGCGC G CAGCCAAU	1394	ATTGGCTG GGCTAGCTACAACGA GCGCCATC	3722
363	GGCGCGCA G CCAAUGGU	1395	ACCATTGG GGCTAGCTACAACGA TGGCGGCC	3723
367	CGCAGCCA A UGGUAGGC	1396	GCCTACCA GGCTAGCTACAACGA TGGCTGCG	3724
370	AGCCAAUG G UAGGCCGC	1397	GCGGCCCTA GGCTAGCTACAACGA CATTGGCT	3725
374	AAUGGUAG G CCGCGCCU	1398	AGGCGCGG GGCTAGCTACAACGA CTACCATT	3726
377	GGUAGGCC G CGCCUGGC	1399	GCCAGGCG GGCTAGCTACAACGA GGCCTACC	3727
379	UAGGCCGC G CCUGGCAG	1400	CTGCCAGG GGCTAGCTACAACGA GCGGCCCTA	3728
384	CGCGCCUG G CAGACGGA	1401	TCCGTCTG GGCTAGCTACAACGA CAGGCGCG	3729
388	CCUGGCAG A CGGACGGG	1402	CCCGTCCG GGCTAGCTACAACGA CTGCCAGG	3730
392	GCAGACGG A CGGGCGCG	1403	CGCGCCCG GGCTAGCTACAACGA CCGTCTGC	3731
396	ACGGACGG G CGCGGGGC	1404	GCCCCGCG GGCTAGCTACAACGA CCGTCCGT	3732
398	GGACGGGC G CGGGGCGG	1405	CCGCCCGG GGCTAGCTACAACGA GCGCGTCC	3733
403	GGCGCGGG G CGGGGCGU	1406	ACGCCCGG GGCTAGCTACAACGA CCCGCGCC	3734
408	GGGGCGGG G CGUGCGCA	1407	TGCGCACG GGCTAGCTACAACGA CCCGCCCC	3735
410	GGCGGGGC G UGCGCAGG	1408	CCTGCGCA GGCTAGCTACAACGA GCGCGGCC	3736
412	CGGGCGGU G CGCAGGCC	1409	GGCCTGCG GGCTAGCTACAACGA ACGCCCCG	3737
414	GGGCGUGC G CAGGCCCCG	1410	CGGGCCTG GGCTAGCTACAACGA GCACGCCC	3738
418	GUGCGCAG G CCCGCCCCG	1411	CGGGCGGG GGCTAGCTACAACGA CTGCGCAC	3739
422	GCAGCCCC G CCCGAGUC	1412	GACTCGGG GGCTAGCTACAACGA GGGCCTGC	3740
428	CCGCCCGA G UCUCGCGC	1413	GGCGGAGA GGCTAGCTACAACGA TCGGGCGG	3741
434	GAGUCUCC G CCGCCCGU	1414	ACGGGCGG GGCTAGCTACAACGA GGAGACTC	3742
437	UCUCGCGC G CCCGUGCC	1415	GGCACGGG GGCTAGCTACAACGA GCGGAGA	3743
441	CGCCGCCC G UGCCCUGC	1416	GCAGGGCA GGCTAGCTACAACGA GGGCGGCG	3744
443	CCGCCCGU G CCCUGCGC	1417	GCGCAGGG GGCTAGCTACAACGA ACGGGCGG	3745
448	CGUGCCCU G CGCCCGCA	1418	TGCGGGCG GGCTAGCTACAACGA AGGGCACG	3746
450	UGCCCUGC G CCCGCAAC	1419	GTTGCGGG GGCTAGCTACAACGA GCAGGGCA	3747
454	CUGCGCCC G CAACCCGA	1420	TCGGGTTG GGCTAGCTACAACGA GGGCGCAG	3748
457	CGCCCGCA A CCCGAGCC	1421	GGCTCGGG GGCTAGCTACAACGA TGGGGCGG	3749

463	CAACCCGA G CCGCACCC	1422	GGGTGCGG GGCTAGCTACAACGA TCGGGTTG	3750
466	CCCGAGCC G CACCCGCC	1423	GGCGGGTG GGCTAGCTACAACGA GGCTCGGG	3751
468	CGAGCCGC A CCGCCGCG	1424	GCGGCGGG GGCTAGCTACAACGA GCGGCTCG	3752
472	CCGCACCC G CCGCGGAC	1425	GTCCGCGG GGCTAGCTACAACGA GGGTGC GG	3753
475	CACCCGCC G CGGACGGA	1426	TCCGTCCG GGCTAGCTACAACGA GGCGGGTG	3754
479	CGCCGCGG A CGGAGCCC	1427	GGGCTCCG GGCTAGCTACAACGA CCGCGGCG	3755
484	CGGACGGA G CCCAUGCG	1428	CGCATGGG GGCTAGCTACAACGA TCCGTCCG	3756
488	CGGAGCCC A UGCGCGGG	1429	CCCCGCGA GGCTAGCTACAACGA GGGCTCCG	3757
490	GAGCCCAU G CGCGGGGC	1430	GCCCCGCG GGCTAGCTACAACGA ATGGGCTC	3758
492	GCCCAUGC G CGGGGCGA	1431	TGCGCCCG GGCTAGCTACAACGA GCATGGGC	3759
497	UGCGCGGG G CGAACCGC	1432	GCGGTTCC GGCTAGCTACAACGA CCGCGCGA	3760
501	CGGGGCGA A CCGCGCGC	1433	GCGCGCGG GGCTAGCTACAACGA TCGCCCCG	3761
504	GGCGAACG G CGCGCCCC	1434	GGGGCGCG GGCTAGCTACAACGA GGTTCCGC	3762
506	CGAACCGC G CGCCCCCG	1435	CGGGGGCG GGCTAGCTACAACGA GCGGTTCC	3763
508	AACCGCGC G CCCCCGCC	1436	GGCGGGGG GGCTAGCTACAACGA GCGCGGTT	3764
514	GCGCCCCC G CCCCCGCC	1437	GGCGGGGG GGCTAGCTACAACGA GGGGGCGC	3765
520	CCGCCCCC G CCGCGCCC	1438	GGGCGGGG GGCTAGCTACAACGA GGGGGCGG	3766
525	CCCGCCCC G CCGCGGCC	1439	GGCGGGGG GGCTAGCTACAACGA GGGGCGGG	3767
531	CCGCCCCG G CCUCGGCC	1440	GGCCGAGG GGCTAGCTACAACGA CGGGGCGG	3768
537	CGGCCUCG G CCGCGGCC	1441	GGCCGGGG GGCTAGCTACAACGA CGAGGCCG	3769
543	CGGCCCCG G CCCUGGCC	1442	GGCCAGGG GGCTAGCTACAACGA CGGGGCCG	3770
549	CGGCCCCG G CCGCGGGG	1443	CCCCGGGG GGCTAGCTACAACGA CAGGGCCG	3771
558	CCCCGGGG G CAGUCGCG	1444	CGCGACTG GGCTAGCTACAACGA CCGCGGGG	3772
561	CGGGGGCA G UCGCGCCU	1445	AGGCGCGA GGCTAGCTACAACGA TGCCCCCG	3773
564	GGGCAGUC G CGCCUGUG	1446	CACAGGCG GGCTAGCTACAACGA GACTGCCC	3774
566	GCAGUCGC G CCUGUGAA	1447	TTCACAGG GGCTAGCTACAACGA GCGACTGC	3775
570	UCGCGCCU G UGAACGGU	1448	ACCGTTCA GGCTAGCTACAACGA AGGCGCGA	3776
574	GCCUGUGA A CGGUGAGU	1449	ACTCACC G GGCTAGCTACAACGA TCACAGGC	3777
577	UGUGAACG G UGAGUGCG	1450	CGCACTCA GGCTAGCTACAACGA CGTTTACA	3778
581	AACGGUGA G UGCGGGCA	1451	TGCCCCGA GGCTAGCTACAACGA TCACCGTT	3779
583	CGGUGAGU G CGGGCAGG	1452	CCTGCCCC GGCTAGCTACAACGA ACTCACC G	3780
587	GAGUGCGG G CAGGGAUC	1453	GATCCCTG GGCTAGCTACAACGA CCGCACTC	3781
593	GGGCAGGG A UCGGCCGG	1454	CCGGCCGA GGCTAGCTACAACGA CCCTGCCC	3782
597	AGGGAUCG G CCGGGCCG	1455	CGGCCCGG GGCTAGCTACAACGA CGATCCCT	3783
602	UCGGCCCG G CCGCGCGC	1456	GCGCGCGG GGCTAGCTACAACGA CCGGCCGA	3784
605	GCCGGGCC G CGCGCCCU	1457	AGGGCGCG GGCTAGCTACAACGA GGCCC GCG	3785
607	CGGGCCGC G CGCCUCC	1458	GGAGGGCG GGCTAGCTACAACGA GCGGCCCG	3786
609	GGCCGCGC G CCCUCCUC	1459	GAGGAGGG GGCTAGCTACAACGA GCGCGGCC	3787
618	CCCUCCUC G CCCCCAGG	1460	CCTGGGGG GGCTAGCTACAACGA GAGGAGGG	3788
626	GCCCCCAG G CGGCAGCA	1461	TGCTGCCG GGCTAGCTACAACGA CTGGGGGC	3789
629	CCCAGGCG G CAGCAUAU	1462	TATTGCTG GGCTAGCTACAACGA CGCCTGGG	3790
632	AGGCGGCA G CAAUACGC	1463	GCGTATTG GGCTAGCTACAACGA TGCCGCCT	3791
635	CGGCAGCA A UACGCGCG	1464	CGCGCGTA GGCTAGCTACAACGA TGCTGCCG	3792
637	GCAGCAAU A CGCGGGC	1465	GCCGCGCG GGCTAGCTACAACGA ATTGCTGC	3793
639	AGCAAUAC G CGCGGCGC	1466	GCGCCGCG GGCTAGCTACAACGA GTATTGCT	3794
641	CAAUACGC G CGGCGCGG	1467	CCGCGCGG GGCTAGCTACAACGA GCGTATTG	3795
644	UACGCGCG G CGCGGGC	1468	GGCCCCGCG GGCTAGCTACAACGA CGCGCGTA	3796
646	CGCGCGGC G CGGGCCGG	1469	CCGGCCCC GGCTAGCTACAACGA GCCGCGCG	3797
650	CGGCGCGG G CCGGGGGC	1470	GCCCCCGG GGCTAGCTACAACGA CCGCGCCG	3798
657	GGCCGGGG G CGCGGGC	1471	GCCCCCGG GGCTAGCTACAACGA CCGCGGCC	3799
659	CGGGGGGC G CGGGCCG	1472	CGGCCCCG GGCTAGCTACAACGA GCCCCCGG	3800
664	GGCGGGG G CCGGCGGG	1473	CCCCCGCG GGCTAGCTACAACGA CCGCGGCC	3801

668	CGGGGCCG G CGGGCGUA	1474	TACGCCCG GGCTAGCTACAACGA CGGCCCCG	3802
672	GCCGGCCG G CGUAAGCG	1475	CGCTTACG GGCTAGCTACAACGA CCGCCGGC	3803
674	CGGCGGGC G UAAGCGGC	1476	GCCGCTTA GGCTAGCTACAACGA GCCCGCCG	3804
678	GGGCGUAA G CGGCGGCG	1477	CGCCGCCG GGCTAGCTACAACGA TTACGCC	3805
681	CGUAAGCG G CGGCGGCG	1478	CGCCGCCG GGCTAGCTACAACGA CGCTTACG	3806
684	AAGCGGCG G CGGCGGCG	1479	CGCCGCCG GGCTAGCTACAACGA CGCCGCTT	3807
687	CGGCGGCG G CGGCGGCG	1480	CGCCGCCG GGCTAGCTACAACGA CGCCGCCG	3808
690	CGGCGGCG G CGGCGGGU	1481	ACCCGCCG GGCTAGCTACAACGA CGCCGCCG	3809
693	CGGCGGCG G CGGGUGGG	1482	CCCACCCG GGCTAGCTACAACGA CGCCGCCG	3810
697	GGCGGCGG G UGGGUGGG	1483	CCCACCCA GGCTAGCTACAACGA CCGCCGCC	3811
701	GCGGGUGG G UGGGGCCG	1484	CGGCCCCA GGCTAGCTACAACGA CCACCCGC	3812
706	UGGGUGGG G CCGGGCGG	1485	CCGCCCGG GGCTAGCTACAACGA CCCACCCA	3813
711	GGGGCCCG G CGGGGCC	1486	GGGCCCCG GGCTAGCTACAACGA CCGGCCCC	3814
716	CGGGCGGG G CCGCGGG	1487	CCCGCGGG GGCTAGCTACAACGA CCCGCCCG	3815
720	CGGGGCCG G CGGGCACA	1488	TGTGCCCC GGCTAGCTACAACGA GGGCCCCG	3816
724	GCCCGCGG G CACAGGUG	1489	CACCTGTG GGCTAGCTACAACGA CCGCGGGC	3817
726	CCGCGGGC A CAGGUGAG	1490	CTCACCTG GGCTAGCTACAACGA GCCCGCGG	3818
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734	ACAGGUGA G CGGGCGUC	1492	GACGCCCC GGCTAGCTACAACGA TCACCTGT	3820
738	GUGAGCGG G CGUCGGGG	1493	CCCCGACG GGCTAGCTACAACGA CCGCTCAC	3821
740	GAGCGGGC G UCGGGGGC	1494	GCCCCCGA GGCTAGCTACAACGA GCCCGCTC	3822
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757	UGCGGGCG G CGGGGGCC	1498	GGCCCCCG GGCTAGCTACAACGA CCGCCGCA	3826
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784	UGGGGCCU G CGGGAAUC	1501	GATTCCCG GGCTAGCTACAACGA AGGCCCCA	3829
790	CUGCGGGA A UCCGGGCC	1502	GGCCCGGA GGCTAGCTACAACGA TCCCGCAG	3830
796	GAAUCCGG G CCCCACCC	1503	GGGTGGGG GGCTAGCTACAACGA CCGGATTC	3831
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805	CCCCACCC G UGGCCUCG	1505	CGAGGCCA GGCTAGCTACAACGA GGGTGGGG	3833
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813	GUGGCCUC G CGCUGGGC	1507	GCCCAGCG GGCTAGCTACAACGA GAGGCCAC	3835
815	GGCCUCGC G CUGGGCAC	1508	GTGCCCAG GGCTAGCTACAACGA GCGAGGCC	3836
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822	CGCUGGGC A CGGUCCCC	1510	GGGGACCG GGCTAGCTACAACGA GCCCAGCG	3838
825	UGGGCACG G UCCCCACG	1511	CGTGGGGA GGCTAGCTACAACGA CGTGCCCA	3839
831	CGGUCCCC A CGCCGGCG	1512	CGCCGGCG GGCTAGCTACAACGA GGGGACCG	3840
833	GUCCCCAC G CCGCGGUA	1513	TACGCCCG GGCTAGCTACAACGA GTGGGGAC	3841
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849	ACCCGGGA G CCUCGGGC	1517	GCCCGAGG GGCTAGCTACAACGA TCCCGGGT	3845
856	AGCCUCGG G CCCGGCGC	1518	GCGCCGGG GGCTAGCTACAACGA CCGAGGCT	3846
861	CGGGCCCG G CGCCCUCA	1519	TGAGGGCG GGCTAGCTACAACGA CGGGCCCC	3847
863	GGCCCGGC G CCCUCACA	1520	TGTGAGGG GGCTAGCTACAACGA GCCGGGCC	3848
869	GCGCCUC A CACCCGGG	1521	CCCGGGTG GGCTAGCTACAACGA GAGGGCGC	3849
871	GCCCUCAC A CCCGGGGG	1522	CCCCGGGG GGCTAGCTACAACGA GTGAGGGC	3850
879	ACCCGGGG G CGUCUGGG	1523	CCCAGACG GGCTAGCTACAACGA CCCCGGGT	3851
881	CCGGGGGC G UCUGGGAG	1524	CTCCCGAG GGCTAGCTACAACGA GCCCGCGG	3852
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905	CCGCGGCC A CGGCACGC	1529	GCGTGCCG GGCTAGCTACAACGA GGCCGCGG	3857
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912	CACGGCAC G CCCGGGCA	1532	TGCCCCGG GGCTAGCTACAACGA GTGCCGTG	3860
918	ACGCCCGG G CACCCCGG	1533	CGGGGGTG GGCTAGCTACAACGA CCGGGCGT	3861
920	CCCCGGGC A CCCCCGAU	1534	ATCGGGGG GGCTAGCTACAACGA GCCCGGGC	3862
927	CACCCCGG A UUCAGCAU	1535	ATGCTGAA GGCTAGCTACAACGA CGGGGGTG	3863
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974	GCCCCAGU G CCUUUUC	1546	GGAAAAGG GGCTAGCTACAACGA ACTGGGGC	3874
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1149	GGCGCCUA G UACGCAGU	1575	ACTGCGTA GGCTAGCTACAACGA TAGGCGCC	3903
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1162	CAGUAGGC G CUCAGCAA	1580	TTGCTGAG GGCTAGCTACAACGA GCCTACTG	3908
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1173	CAGCAAU A CUUGUCGG	1583	CCGACAAG GGCTAGCTACAACGA ATTTGCTG	3911
1177	AAUACUU G UCGGAGGC	1584	GCCTCCGA GGCTAGCTACAACGA AAGTATTT	3912
1184	UGUCGGAG G CACCAGCG	1585	CGCTGGTG GGCTAGCTACAACGA CTCCGACA	3913
1186	UCGGAGGC A CCAGCGCC	1586	GGCGCTGG GGCTAGCTACAACGA GCCTCCGA	3914
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1192	GCACCAGC G CCGCGGG	1588	CCCCGCGG GGCTAGCTACAACGA GCTGGTGC	3916
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1332	AGCUUCUA A UUUGGGUG	1620	CACCCAAA GGCTAGCTACAACGA TAGAAGCT	3948
1338	UAAUUUGG G UGCGUGGU	1621	ACCACGCA GGCTAGCTACAACGA CCAAATTA	3949
1340	AUUUGGGU G CGUGGUUG	1622	CAACCACG GGCTAGCTACAACGA ACCCAAAT	3950
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1345	GGUGCGUG G UUGAGAGC	1624	GCTCTCAA GGCTAGCTACAACGA CACGCACC	3952
1352	GGUUGAGA G CGCUCAGC	1625	GCTGAGCG GGCTAGCTACAACGA TCTCAACC	3953
1354	UUGAGAGC G CUCAGCUG	1626	CAGCTGAG GGCTAGCTACAACGA GCTCTCAA	3954
1359	AGCGCUCA G CUGUCAGC	1627	GCTGACAG GGCTAGCTACAACGA TGAGCGCT	3955
1362	GCUCAGCU G UCAGCCCU	1628	AGGGCTGA GGCTAGCTACAACGA AGCTGAGC	3956
1366	AGCUGUCA G CCCUGCCU	1629	AGGCAGGG GGCTAGCTACAACGA TGACAGCT	3957

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1386	AGGGCUGG G UCCCUUUU	1632	AAAAGGGA GGCTAGCTACAACGA CCAGCCCT	3960
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1497	GCUGCCUG G CGUUGGGG	1659	CCCCAACG GGCTAGCTACAACGA CAGGCAGC	3987
1499	UGCCUGGC G UUGGGGCC	1660	GGCCCCAA GGCTAGCTACAACGA GCCAGGCA	3988
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1513	GCCCAGGG A CCGCUGUG	1662	CACAGCGG GGCTAGCTACAACGA CCCTGGGC	3990
1516	CAGGGACC G CUGUGGGU	1663	ACCCACAG GGCTAGCTACAACGA GGTCCCTG	3991
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1527	GUGGGUUU G CCCUUCAG	1666	CTGAAGGG GGCTAGCTACAACGA AAACCCAC	3994
1536	CCCUUCAG A UGCCCUG	1667	CAGGGCCA GGCTAGCTACAACGA CTGAAGGG	3995
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1544	AUGGCCCC G CCAGCAGC	1669	GCTGCTGG GGCTAGCTACAACGA AGGGCCAT	3997
1548	CCUGGCCA G CAGCUGCC	1670	GGCAGCTG GGCTAGCTACAACGA TGGCAGGG	3998
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1554	CAGCAGCU G CCCUGGG	1672	CCACAGGG GGCTAGCTACAACGA AGCTGCTG	4000
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1582	GGGCCUGG G CCUGGCUG	1677	CAGCCAGG GGCTAGCTACAACGA CCAGGCCC	4005
1587	UGGGCCUG G CUGAGCAG	1678	CTGCTCAG GGCTAGCTACAACGA CAGGCCCA	4006
1592	CUGGCUGA G CAGGGCCC	1679	GGGCCCTG GGCTAGCTACAACGA TCAGCCAG	4007
1597	UGAGCAGG G CCCUCCUU	1680	AAGGAGGG GGCTAGCTACAACGA CCTGCTCA	4008
1607	CCUCCUUG G CAGGUGGG	1681	CCCACCTG GGCTAGCTACAACGA CAAGGAGG	4009

1611	CUUGGCAG G UGGGGCAG	1682	CTGCCCCA GGCTAGCTACAACGA CTGCCAAG	4010
1616	CAGGUGGG G CAGGAGAC	1683	GTCTCCTG GGCTAGCTACAACGA CCCACCTG	4011
1623	GGCAGGAG A CCCUGUAG	1684	CTACAGGG GGCTAGCTACAACGA CTCCTGCC	4012
1628	GAGACCCU G UAGGAGGA	1685	TCCTCCTA GGCTAGCTACAACGA AGGGTCTC	4013
1636	GUAGGAGG A CCCCGGGC	1686	GCCCGGGG GGCTAGCTACAACGA CCTCTAC	4014
1643	GACCCCGG G CCGCAGGC	1687	GCCTGCCG GGCTAGCTACAACGA CCGGGGTC	4015
1646	CCCGGGCC G CAGGCCCC	1688	GGGGCCTG GGCTAGCTACAACGA GGCCCGGG	4016
1650	GGCCGCAG G CCCUGAG	1689	CTCAGGGG GGCTAGCTACAACGA CTGCGGCC	4017
1661	CCUGAGGA G CGAUGACG	1690	CGTCATCG GGCTAGCTACAACGA TCCTCAGG	4018
1664	GAGGAGCG A UGACGGAA	1691	TTCCGTCA GGCTAGCTACAACGA CGCTCCTC	4019
1667	GAGCGAUG A CGGAUAU	1692	ATATTCGG GGCTAGCTACAACGA CATCGCTC	4020
1672	AUGACGGA A UAUAAGCU	1693	AGCTTATA GGCTAGCTACAACGA TCCGTCTC	4021
1674	GACGGAAU A UAAGCUGG	1694	CCAGCTTA GGCTAGCTACAACGA ATTCCGTC	4022
1678	GAAUAUAA G CUGGUGGU	1695	ACCACCAG GGCTAGCTACAACGA TTATATTC	4023
1682	AUAAGCUG G UGGUGGUG	1696	CACCACCA GGCTAGCTACAACGA CAGCTTAT	4024
1685	AGCUGGUG G UGGUGGGC	1697	GCCCACCA GGCTAGCTACAACGA CACCAGCT	4025
1688	UGGUGGUG G UGGGCGCC	1698	GGCGCCCA GGCTAGCTACAACGA CACCACCA	4026
1692	GGUGGUGG G CGCCGGCG	1699	CGCCGGCG GGCTAGCTACAACGA CCACCACC	4027
1694	UGGUGGGC G CCGGCGGU	1700	ACCGCCGG GGCTAGCTACAACGA GCCCACC	4028
1698	GGGCGCCG G CGGUGUGG	1701	CCACACCG GGCTAGCTACAACGA CGGCGCCC	4029
1701	CGCCGGCG G UGUGGGCA	1702	TGCCCCA GGCTAGCTACAACGA CGCCGGCG	4030
1703	CCGGCGGU G UGGGCAAG	1703	CTTGCCCC GGCTAGCTACAACGA ACCGCCGG	4031
1707	CGGUGUGG G CAAGAGUG	1704	CACTCTTG GGCTAGCTACAACGA CCACACCG	4032
1713	GGGCAAGA G UGCGUGA	1705	TCAGCGCA GGCTAGCTACAACGA TCTTGCCC	4033
1715	GCAAGAGU G CGCUGACC	1706	GGTCAGCG GGCTAGCTACAACGA ACTCTTGC	4034
1717	AAGAGUGC G CUGACCAU	1707	ATGGTCAG GGCTAGCTACAACGA GCACTCTT	4035
1721	GUGCGCUG A CCAUCCAG	1708	CTGGATGG GGCTAGCTACAACGA CAGCGCAC	4036
1724	CGCUGACC A UCCAGCUG	1709	CAGCTGGA GGCTAGCTACAACGA GGTCAGCG	4037
1729	ACCAUCCA G CUGAUCCA	1710	TGGATCAG GGCTAGCTACAACGA TGGATGGT	4038
1733	UCCAGCUG A UCCAGAAC	1711	GTTCTGGA GGCTAGCTACAACGA CAGCTGGA	4039
1740	GAUCCAGA A CCAUUUUG	1712	CAAAATGG GGCTAGCTACAACGA TCTGGATC	4040
1743	CCAGAACC A UUUUGUGG	1713	CCACAAAA GGCTAGCTACAACGA GGTTCTGG	4041
1748	ACCAUUUU G UGGACGAA	1714	TTCGTCCA GGCTAGCTACAACGA AAAATGGT	4042
1752	UUUUGUGG A CGAAUACG	1715	CGTATTCG GGCTAGCTACAACGA CCACAAAA	4043
1756	GUGGACGA A UACGACCC	1716	GGGTCGTA GGCTAGCTACAACGA TCGTCCAC	4044
1758	GGACGAAU A CGACCCCA	1717	TGGGGTCG GGCTAGCTACAACGA ATTCTGTC	4045
1761	CGAAUACG A CCCACUA	1718	TAGTGGGG GGCTAGCTACAACGA CGTATTCG	4046
1766	ACGACCCC A CUAUAGAG	1719	CTCTATAG GGCTAGCTACAACGA GGGGTCGT	4047
1769	ACCCACU A UAGAGGAU	1720	ATCCTCTA GGCTAGCTACAACGA AGTGGGGT	4048
1776	UAUAGAGG A UUCUACC	1721	GGTAGGAA GGCTAGCTACAACGA CCTCTATA	4049
1782	GGAUUCCU A CCGGAAGC	1722	GCTTCCGG GGCTAGCTACAACGA AGGAATCC	4050
1789	UACCGGAA G CAGGUGGU	1723	ACCACCTG GGCTAGCTACAACGA TTCCGGTA	4051
1793	GGAAGCAG G UGGUCAU	1724	AATGACCA GGCTAGCTACAACGA CTGCTTCC	4052
1796	AGCAGGUG G UCAUUGAU	1725	ATCAATGA GGCTAGCTACAACGA CACCTGCT	4053
1799	AGGUGGUC A UUGAUGGG	1726	CCCATCAA GGCTAGCTACAACGA GACCACCT	4054
1803	GGUCAUUG A UGGGGAGA	1727	TCTCCCCA GGCTAGCTACAACGA CAATGACC	4055
1811	AUGGGGAG A CGUGCCUG	1728	CAGGCACG GGCTAGCTACAACGA CTCCCCAT	4056
1813	GGGGAGAC G UGCCUGUU	1729	AACAGGCA GGCTAGCTACAACGA GTCTCCCC	4057
1815	GGAGACGU G CCUGUUGG	1730	CCAACAGG GGCTAGCTACAACGA ACGTCTCC	4058
1819	ACGUGCCU G UUGGACAU	1731	ATGTCCAA GGCTAGCTACAACGA AGGCACGT	4059
1824	CCUGUUGG A CAUCCUGG	1732	CCAGGATG GGCTAGCTACAACGA CCAACAGG	4060
1826	UGUUGGAC A UCCUGGAU	1733	ATCCAGGA GGCTAGCTACAACGA GTCCAACA	4061

1833	CAUCCUGG A UACCGCCG	1734	CGGCGGTA GGCTAGCTACAACGA CCAGGATG	4062
1835	UCCUGGAU A CCGCCGGC	1735	GCCGGCGG GGCTAGCTACAACGA ATCCAGGA	4063
1838	UGGAUACC G CCGGCCAG	1736	CTGGCCGG GGCTAGCTACAACGA GGTATCCA	4064
1842	UACCGCCG G CCAGGAGG	1737	CCTCCTGG GGCTAGCTACAACGA CGGCGGTA	4065
1852	CAGGAGGA G UACAGCGC	1738	GCGCTGTA GGCTAGCTACAACGA TCCTCCTG	4066
1854	GGAGGAGU A CAGCGCCA	1739	TGGCGCTG GGCTAGCTACAACGA ACTCCTCC	4067
1857	GGAGUACA G CGCCAUGC	1740	GCATGGCG GGCTAGCTACAACGA TGTACTCC	4068
1859	AGUACAGC G CCAUGCGG	1741	CCGCATGG GGCTAGCTACAACGA GCTGTACT	4069
1862	ACAGCGCC A UGCGGGAC	1742	GTCCCCGA GGCTAGCTACAACGA GCGCTGT	4070
1864	AGCGCCAU G CGGGACCA	1743	TGGTCCCG GGCTAGCTACAACGA ATGGCGCT	4071
1869	CAUGCGGG A CCAGUACA	1744	TGTACTGG GGCTAGCTACAACGA CCGCATG	4072
1873	CGGGACCA G UACAUGC	1745	CGCATGTA GGCTAGCTACAACGA TGGTCCCG	4073
1875	GGACCAGU A CAUGCGCA	1746	TGCGCATG GGCTAGCTACAACGA ACTGGTCC	4074
1877	ACCAGUAC A UGCGCACC	1747	GGTGCACA GGCTAGCTACAACGA GTACTGGT	4075
1879	CAGUACAU G CGCACCAG	1748	CCGGTGCG GGCTAGCTACAACGA ATGTACTG	4076
1881	GUACAUGC G CACCGGGG	1749	CCCCGGTG GGCTAGCTACAACGA GCATGTAC	4077
1883	ACAUGC GC A CCGGGGAG	1750	CTCCCCGG GGCTAGCTACAACGA GCGCATGT	4078
1893	CGGGGAGG G CUUCCUGU	1751	ACAGGAAG GGCTAGCTACAACGA CCTCCCCG	4079
1900	GGCUUCCU G UGUGUGUU	1752	AACACACA GGCTAGCTACAACGA AGGAAGCC	4080
1902	CUUCCUGU G UGUGUUUG	1753	CAAACACA GGCTAGCTACAACGA ACAGGAAG	4081
1904	UCCUGUGU G UGUUUGCC	1754	GGCAAACA GGCTAGCTACAACGA ACACAGGA	4082
1906	CUGUGUGU G UUUGCCAU	1755	ATGGCAAA GGCTAGCTACAACGA ACACACAG	4083
1910	GUGUGUUU G CCAUCAAC	1756	GTTGATGG GGCTAGCTACAACGA AAACACAC	4084
1913	UGUUUGCC A UCAACAAC	1757	GTTGTTGA GGCTAGCTACAACGA GGCAAACA	4085
1917	UGCCAUCA A CAACACCA	1758	TGGTGTGG GGCTAGCTACAACGA TGATGGCA	4086
1920	CAUCAACA A CACCAAGU	1759	ACTTGGTG GGCTAGCTACAACGA TGTGTATG	4087
1922	UCAACAAC A CCAAGUCU	1760	AGACTTGG GGCTAGCTACAACGA GTTGTGTA	4088
1927	AACACCAA G UCUUUUGA	1761	TCAAAAGA GGCTAGCTACAACGA TTGGTGTT	4089
1938	UUUUGAGG A CAUCCACC	1762	GGTGGATG GGCTAGCTACAACGA CCTCAAAA	4090
1940	UUGAGGAC A UCCACCAG	1763	CTGGTGGA GGCTAGCTACAACGA GTCCTCAA	4091
1944	GGACAUCC A CCAGUACA	1764	TGTACTGG GGCTAGCTACAACGA GGATGTCC	4092
1948	AUCCACCA G UACAGGGA	1765	TCCCTGTA GGCTAGCTACAACGA TGGTGGAT	4093
1950	CCACCAGU A CAGGGAGC	1766	GCTCCCTG GGCTAGCTACAACGA ACTGGTGG	4094
1957	UACAGGGA G CAGAUCAA	1767	TTGATCTG GGCTAGCTACAACGA TCCCTGTA	4095
1961	GGGAGCAG A UCAAACGG	1768	CCGTTTGA GGCTAGCTACAACGA CTGCTCCC	4096
1966	CAGAUCAA A CGGGUGAA	1769	TTCACCCG GGCTAGCTACAACGA TTGATCTG	4097
1970	UCAAACGG G UGAAGGAC	1770	GTCCTTCA GGCTAGCTACAACGA CCGTTTGA	4098
1977	GGUGAAGG A CUCGGAUG	1771	CATCCGAG GGCTAGCTACAACGA CCTTCACC	4099
1983	GGACUCGG A UGACGUGC	1772	GCACGTCA GGCTAGCTACAACGA CCGAGTCC	4100
1986	CUCGGAUG A CGUGCCCA	1773	TGGGCACG GGCTAGCTACAACGA CATCCGAG	4101
1988	CGGAUGAC G UGCCCAUG	1774	CATGGGCA GGCTAGCTACAACGA GTCATCCG	4102
1990	GAUGACGU G CCCAUGGU	1775	ACCATGGG GGCTAGCTACAACGA ACGTCATC	4103
1994	ACGUGCCC A UGGUGCUG	1776	CAGCACCA GGCTAGCTACAACGA GGGCACGT	4104
1997	UGCCCAUG G UGCUGGUG	1777	CACCAGCA GGCTAGCTACAACGA CATGGGCA	4105
1999	CCCAUGGU G CUGGUGGG	1778	CCCACCAG GGCTAGCTACAACGA ACCATGGG	4106
2003	UGGUGCUG G UGGGGAAC	1779	GTTCCCCA GGCTAGCTACAACGA CAGCACCA	4107
2010	GGUGGGGA A CAAGUGUG	1780	CACACTTG GGCTAGCTACAACGA TCCCCACC	4108
2014	GGGAACAA G UGUGACCU	1781	AGGTCACA GGCTAGCTACAACGA TTGTTCCC	4109
2016	GAACAAGU G UGACCUGG	1782	CCAGGTCA GGCTAGCTACAACGA ACTTGTTT	4110
2019	CAAGUGUG A CCUGGCUG	1783	CAGCCAGG GGCTAGCTACAACGA CACACTTG	4111
2024	GUGACCUG G CUGCACGC	1784	GCGTGCAG GGCTAGCTACAACGA CAGGTCAC	4112
2027	ACCUGGCU G CACGCACU	1785	AGTGCGTG GGCTAGCTACAACGA AGCCAGGT	4113

2029	CUGGUGC A CGCACUGU	1786	ACAGTGCG GGCTAGCTACAACGA GCAGCCAG	4114
2031	GGCUGCAC G CACUGUGG	1787	CCACAGTG GGCTAGCTACAACGA GTGCAGCC	4115
2033	CUGCACGC A CUGUGGAA	1788	TTCCACAG GGCTAGCTACAACGA GCGTGCAG	4116
2036	CACGCACU G UGGAUUCU	1789	AGATTCCA GGCTAGCTACAACGA AGTGCGTG	4117
2041	ACUGUGGA A UCUCGGCA	1790	TGCCGAGA GGCTAGCTACAACGA TCCACAGT	4118
2047	GAAUCUCG G CAGGCUCA	1791	TGAGCCTG GGCTAGCTACAACGA CGAGATTC	4119
2051	CUCGGCAG G CUCAGGAC	1792	GTCCTGAG GGCTAGCTACAACGA CTGCCGAG	4120
2058	GGCUCAGG A CCUCGCCC	1793	GGCGCAGG GGCTAGCTACAACGA CCTGAGCC	4121
2063	AGGACCUC G CCCGAAGC	1794	GCTTCGGG GGCTAGCTACAACGA GAGGTCCT	4122
2070	CGCCCGAA G CUACGGCA	1795	TGCCGTAG GGCTAGCTACAACGA TTCGGGCG	4123
2073	CCGAAGCU A CGGCAUCC	1796	GGATGCCG GGCTAGCTACAACGA AGCTTCGG	4124
2076	AAGCUACG G CAUCCCUU	1797	AGGGGATG GGCTAGCTACAACGA CGTAGCTT	4125
2078	GPUACGGC A UCCCUUAC	1798	GTAGGGGA GGCTAGCTACAACGA GCCGTAGC	4126
2085	CAUCCCUU A CAUCGAGA	1799	TCTCGATG GGCTAGCTACAACGA AGGGGATG	4127
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2093	ACAUCGAG A CCUCGGCC	1801	GGCCGAGG GGCTAGCTACAACGA CTCGATGT	4129
2099	AGACCUCG G CCAAGACC	1802	GGTCTTGG GGCTAGCTACAACGA CGAGGTCT	4130
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2110	AAGACCCG G CAGGGAGU	1804	ACTCCCTG GGCTAGCTACAACGA CGGGTCTT	4132
2117	GGCAGGGA G UGGAGGAU	1805	ATCCTCCA GGCTAGCTACAACGA TCCCTGCC	4133
2124	AGUGGAGG A UGCCUUCU	1806	AGAAGGCA GGCTAGCTACAACGA CCTCCACT	4134
2126	UGGAGGAU G CCUUCUAC	1807	GTAGAAGG GGCTAGCTACAACGA ATCCTCCA	4135
2133	UGCCUUCU A CACGUUGG	1808	CCAACGTG GGCTAGCTACAACGA AGAAGGCA	4136
2135	CCUUCUAC A CGUUGGUG	1809	CACCAACG GGCTAGCTACAACGA GTAGAAGG	4137
2137	UUCUACAC G UUGGUGCG	1810	CGCACCAA GGCTAGCTACAACGA GTGTAGAA	4138
2141	ACACGUUG G UGCGUGAG	1811	CTCACGCA GGCTAGCTACAACGA CAACGTGT	4139
2143	ACGUUGGU G CGUGAGAU	1812	ATCTCACG GGCTAGCTACAACGA ACCAACGT	4140
2145	GUUGGUGC G UGAGAUCC	1813	GGATCTCA GGCTAGCTACAACGA GCACCAAC	4141
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2155	GAGAUCCG G CAGCACAA	1815	TTGTGCTG GGCTAGCTACAACGA CGGATCTC	4143
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2196	UGAGAGUG G CCCCAGCU	1824	AGCCGGGG GGCTAGCTACAACGA CACTCTCA	4152
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2205	CCCCAGCU G CAUGAGCU	1826	AGCTCATG GGCTAGCTACAACGA AGCCGGGG	4154
2207	CCGGCUGC A UGAGCUGC	1827	GCAGCTCA GGCTAGCTACAACGA GCAGCCGG	4155
2211	CUGCAUGA G CUGCAAGU	1828	ACTTGCAG GGCTAGCTACAACGA TCATGCAG	4156
2214	CAUGAGCU G CAAGUGUG	1829	CACACTTG GGCTAGCTACAACGA AGCTCATG	4157
2218	AGCUGCAA G UGUGUCU	1830	AGCACACA GGCTAGCTACAACGA TTGCAGCT	4158
2220	CUGCAAGU G UGUCUCU	1831	AGAGCACA GGCTAGCTACAACGA ACTTGCAG	4159
2222	GCAAGUGU G UGCUCUCC	1832	GGAGAGCA GGCTAGCTACAACGA ACACTTGC	4160
2224	AAGUGUGU G CUCUCCUG	1833	CAGGAGAG GGCTAGCTACAACGA ACACACTT	4161
2233	CUCUCCUG A CGCAGGUG	1834	CACCTGCG GGCTAGCTACAACGA CAGGAGAG	4162
2235	CUCCUGAC G CAGGUGAG	1835	CTCACCTG GGCTAGCTACAACGA GTCAGGAG	4163
2239	UGACGCAG G UGAGGGGG	1836	CCCCCTCA GGCTAGCTACAACGA CTGCGTCA	4164
2248	UGAGGGGG A CUCCAGG	1837	CCTGGGAG GGCTAGCTACAACGA CCCCCTCA	4165

2257	CUCCAGG G	CGGCCGCC	1838	GGCGGCCG	GGCTAGCTACAACGA	CCTGGGAG	4166
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2268	GCCGCCAC G	CCCACCGG	1842	CCGGTGGG	GGCTAGCTACAACGA	GTGGCGGC	4170
2272	CCACGCCC A	COGAUGA	1843	TCATCCGG	GGCTAGCTACAACGA	GGGCGTGG	4171
2277	CCCACCGG A	UGACCCCG	1844	CGGGGTCA	GGCTAGCTACAACGA	CCGGTGGG	4172
2280	ACCGGAUG A	CCCCGGCU	1845	AGCCGGGG	GGCTAGCTACAACGA	CATCCGGT	4173
2286	UGACCCCG G	CUCCCCGC	1846	GCGGGGAG	GGCTAGCTACAACGA	CGGGGTCA	4174
2293	GGCUCCCC G	CCCCUGCC	1847	GGCAGGGG	GGCTAGCTACAACGA	GGGGAGCC	4175
2299	CCGCCCCU G	CCGGUCUC	1848	GAGACCGG	GGCTAGCTACAACGA	AGGGGCGG	4176
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2336	UCCCUUGU G	CCCCGCCC	1856	GGGCGGGG	GGCTAGCTACAACGA	ACAAGGGA	4184
2341	UGUGCCCC G	CCCAGCAC	1857	GTGCTGGG	GGCTAGCTACAACGA	GGGGCACA	4185
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2359	AGCUCAGG A	CAUGGAGG	1861	CCTCCATG	GGCTAGCTACAACGA	CCTGAGCT	4189
2361	CUCAGGAC A	UGGAGGUG	1862	CACCTCCA	GGCTAGCTACAACGA	GTCTGTAG	4190
2367	ACAUGGAG G	UGCCGGAU	1863	ATCCGGCA	GGCTAGCTACAACGA	CTCCATGT	4191
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2456	GCCCCAGU A	CCCCGGGA	1876	TCCCCGGG	GGCTAGCTACAACGA	GA CTGGGC	4204
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2480	CCGAGGUG A	CUGCAGAC	1881	GTCTGCAG	GGCTAGCTACAACGA	CACCTCGG	4209
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2487	GACUGCAG A	CCCUCCCA	1883	TGGGAGGG	GGCTAGCTACAACGA	CTGCAGTC	4211
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2504	GGGAGGCU G	UGCACAGA	1885	TCTGTGCA	GGCTAGCTACAACGA	AGCCTCCC	4213
2506	GAGGUGU G	CACAGACU	1886	AGTCTGTG	GGCTAGCTACAACGA	ACAGCCTC	4214
2508	GGCUGUGC A	CAGACUGU	1887	ACAGTCTG	GGCTAGCTACAACGA	GCACAGCC	4215
2512	GUGCACAG A	CUGUCUUG	1888	CAAGACAG	GGCTAGCTACAACGA	CTGTGCAC	4216
2515	CACAGACU G	UCUUGAAC	1889	GTTCAAGA	GGCTAGCTACAACGA	AGTCTGTG	4217

2522	UGUCUUGA A CAUCCCAA	1890	TTGGGATG GGCTAGCTACAACGA TCAAGACA	4218
2524	UCUUGAAC A UCCCAAU	1891	ATTTGGGA GGCTAGCTACAACGA GTTCAAGA	4219
2531	CAUCCCAA A UGCCACCG	1892	CGGTGGCA GGCTAGCTACAACGA TTGGGATG	4220
2533	UCCCAAU G CCACCGGA	1893	TCCGGTGG GGCTAGCTACAACGA ATTTGGGA	4221
2536	CAAUGCC A CCGGAACC	1894	GGTTCCGG GGCTAGCTACAACGA GGCATTTG	4222
2542	CCACCGGA A CCCAGCC	1895	GGCTGGGG GGCTAGCTACAACGA TCCGGTGG	4223
2548	GAACCCCA G CCCUAGC	1896	GCTAAGGG GGCTAGCTACAACGA TGGGGTTC	4224
2555	AGCCCUUA G CUCCCCUC	1897	GAGGGGAG GGCTAGCTACAACGA TAAGGGCT	4225
2568	CCUCCCAG G CCUCUGUG	1898	CACAGAGG GGCTAGCTACAACGA CTGGGAGG	4226
2574	AGGCCUCU G UGGGCCCU	1899	AGGGCCCA GGCTAGCTACAACGA AGAGGCCT	4227
2578	CUCUGUGG G CCCUUGUC	1900	GACAAGGG GGCTAGCTACAACGA CCACAGAG	4228
2584	GGGCCCUU G UCGGGCAC	1901	GTGCCCGA GGCTAGCTACAACGA AAGGGCCC	4229
2589	CUUGUCGG G CACAGAUG	1902	CATCTGTG GGCTAGCTACAACGA CCGACAAG	4230
2591	UGUCGGGC A CAGAUGGG	1903	CCCATCTG GGCTAGCTACAACGA GCCCGACA	4231
2595	GGGCACAG A UGGGAUCA	1904	TGATCCCA GGCTAGCTACAACGA CTGTGCCC	4232
2600	CAGAUGGG A UCACAGUA	1905	TACTGTGA GGCTAGCTACAACGA CCCATCTG	4233
2603	AUGGGAUC A CAGUAAU	1906	ATTTACTG GGCTAGCTACAACGA GATCCCAT	4234
2606	GGAUCACA G UAAUUAU	1907	ATAATTTA GGCTAGCTACAACGA TGTGATCC	4235
2610	CACAGUAA A UUAUUGGA	1908	TCCAATAA GGCTAGCTACAACGA TTACTGTG	4236
2613	AGUAAAUU A UUGGAUGG	1909	CCATCCAA GGCTAGCTACAACGA AATTTACT	4237
2618	AUUAUUGG A UGGUCUUG	1910	CAAGACCA GGCTAGCTACAACGA CCAATAAT	4238
2621	AUUGGAUG G UCUUGAUC	1911	GATCAAGA GGCTAGCTACAACGA CATCCAAT	4239
2627	UGGUCUUG A UCUUGGUU	1912	AACCAAGA GGCTAGCTACAACGA CAAGACCA	4240
2633	UGAUCUUG G UUUUCGGC	1913	GCCGAAAA GGCTAGCTACAACGA CAAGATCA	4241
2640	GGUUUUCG G CUGAGGGU	1914	ACCCTCAG GGCTAGCTACAACGA CGAAAACC	4242
2647	GGCUGAGG G UGGGACAC	1915	GTGTCCCA GGCTAGCTACAACGA CCTCAGCC	4243
2652	AGGGUGGG A CACGUGGC	1916	GCACCGTG GGCTAGCTACAACGA CCCACCCT	4244
2654	GGUGGGAC A CGGUGCGC	1917	GCGCACCG GGCTAGCTACAACGA GTCCCACC	4245
2657	GGGACACG G UGCGGUG	1918	CACGCGCA GGCTAGCTACAACGA CGTGTCCC	4246
2659	GACACGGU G CGCGUGUG	1919	CACACGCG GGCTAGCTACAACGA ACCGTGTC	4247
2661	CACGUGGC G CGUGUGGC	1920	GCCACACG GGCTAGCTACAACGA GCACCGTG	4248
2663	CGGUGCGC G UGUGGCCU	1921	AGGCCACA GGCTAGCTACAACGA GCGCACCG	4249
2665	GUGCGCGU G UGGCCUGG	1922	CCAGGCCA GGCTAGCTACAACGA ACGCGCAC	4250
2668	CGCGUGUG G CCUGGCAU	1923	ATGCCAGG GGCTAGCTACAACGA CACACGCG	4251
2673	GUGGCCUG G CAUGAGGU	1924	ACCTCATG GGCTAGCTACAACGA CAGGCCAC	4252
2675	GGCCUGGC A UGAGGUUA	1925	ATACCTCA GGCTAGCTACAACGA GCCAGGCC	4253
2680	GGCAUGAG G UAUGUCGG	1926	CCGACATA GGCTAGCTACAACGA CTCATGCC	4254
2682	CAUGAGGU A UGUCGGA	1927	TTCCGACA GGCTAGCTACAACGA ACCTCATG	4255
2684	UGAGGUUA G UCGGAACC	1928	GGTTCCGA GGCTAGCTACAACGA ATACCTCA	4256
2690	AUGUCGGA A CCUCAGGC	1929	GCCTGAGG GGCTAGCTACAACGA TCCGACAT	4257
2697	AACCUAG G CCUGUCCA	1930	TGGACAGG GGCTAGCTACAACGA CTGAGGTT	4258
2701	UCAGGCCU G UCCAGCCC	1931	GGGCTGGA GGCTAGCTACAACGA AGGCCTGA	4259
2706	CCUGUCCA G CCCUGGGC	1932	GCCCAGGG GGCTAGCTACAACGA TGGACAGG	4260
2713	AGCCUGG G CUCUCCAU	1933	ATGGAGAG GGCTAGCTACAACGA CCAGGGCT	4261
2720	GGCUCUCC A UAGCCUUU	1934	AAAGGCTA GGCTAGCTACAACGA GGAGAGCC	4262
2723	UCUCCAUA G CCUUUGGG	1935	CCCAAAGG GGCTAGCTACAACGA TATGGAGA	4263
2740	AGGGGGAG G UUGGGAGA	1936	TCTCCCAA GGCTAGCTACAACGA CTCCCCCT	4264
2750	UGGGAGAG G CCGGUCAG	1937	CTGACCGG GGCTAGCTACAACGA CTCTCCCA	4265
2754	AGAGGCCG G UCAGGGGU	1938	ACCCCTGA GGCTAGCTACAACGA CGGCCTCT	4266
2761	GGUCAGGG G UCUGGGCU	1939	AGCCCAGA GGCTAGCTACAACGA CCCTGACC	4267
2767	GGGUCUGG G CUGUGGUG	1940	CACCACAG GGCTAGCTACAACGA CCAGACCC	4268
2770	UCUGGGCU G UGGUGCUC	1941	GAGCACCA GGCTAGCTACAACGA AGCCCAGA	4269

2773	GGGUGUG G UGCUCUCU	1942	AGAGAGCA GGCTAGCTACAACGA CACAGCCC	4270
2775	GCUGUGGU G CUCUCUCC	1943	GGAGAGAG GGCTAGCTACAACGA ACCACAGC	4271
2788	CUCCUCCC G CCUGCCCC	1944	GGGGCAGG GGCTAGCTACAACGA GGGAGGAG	4272
2792	UCCCGCCU G CCCAGUG	1945	CACTGGGG GGCTAGCTACAACGA AGGCGGGA	4273
2798	CUGCCCCA G UGUCCACG	1946	CGTGGACA GGCTAGCTACAACGA TGGGGCAG	4274
2800	GCCCCAGU G UCCACGGC	1947	GCCGTGGA GGCTAGCTACAACGA ACTGGGGC	4275
2804	CAGUGUCC A CGGCUUCU	1948	AGAAGCCG GGCTAGCTACAACGA GGACACTG	4276
2807	UGUCCACG G CUUCUGGC	1949	GCCAGAAG GGCTAGCTACAACGA CGTGGACA	4277
2814	GGCUUCUG G CAGAGAGC	1950	GCTCTCTG GGCTAGCTACAACGA CAGAAGCC	4278
2821	GGCAGAGA G CUCUGGAC	1951	GTCCAGAG GGCTAGCTACAACGA TCTCTGCC	4279
2828	AGCUCUGG A CAAGCAGG	1952	CCTGCTTG GGCTAGCTACAACGA CCAGAGCT	4280
2832	CUGGACAA G CAGGCAGA	1953	TCTGCCTG GGCTAGCTACAACGA TTGTCCAG	4281
2836	ACAAGCAG G CAGAUCAU	1954	ATGATCTG GGCTAGCTACAACGA CTGCTTGT	4282
2840	GCAGGCAG A UCAUAAGG	1955	CCTTATGA GGCTAGCTACAACGA CTGCCTGC	4283
2843	GGCAGAU A UAAGGACA	1956	TGTCCCTTA GGCTAGCTACAACGA GATCTGCC	4284
2849	UCAUAAGG A CAGAGAGC	1957	GCTCTCTG GGCTAGCTACAACGA CCTTATGA	4285
2856	GACAGAGA G CUUACUGU	1958	ACAGTAAG GGCTAGCTACAACGA TCTCTGTC	4286
2860	GAGAGCUU A CUGUGCUU	1959	AAGCACAG GGCTAGCTACAACGA AAGCTCTC	4287
2863	AGCUUACU G UGUUCUA	1960	TAGAAGCA GGCTAGCTACAACGA AGTAAGCT	4288
2865	CUUACUGU G CUUCUACC	1961	GGTAGAAG GGCTAGCTACAACGA ACAGTAAG	4289
2871	GUGCUUCU A CCAACUAG	1962	CTAGTTGG GGCTAGCTACAACGA AGAAGCAC	4290
2875	UUCUACCA A CUAGGAGG	1963	CCTCCTAG GGCTAGCTACAACGA TGGTAGAA	4291
2884	CUAGGAGG G CGUCCUGG	1964	CCAGGACG GGCTAGCTACAACGA CCTCCTAG	4292
2886	AGGAGGGC G UCCUGGUC	1965	GACCAGGA GGCTAGCTACAACGA GCCCTCCT	4293
2892	GCGUCCUG G UCCUCCAG	1966	CTGGAGGA GGCTAGCTACAACGA CAGGACGC	4294
2907	AGAGGGAG G UGUUUUCA	1967	TGAAACCA GGCTAGCTACAACGA CTCCCTCT	4295
2910	GGGAGGUG G UUCAGGG	1968	CCCTGAAA GGCTAGCTACAACGA CACCTCCC	4296
2919	UUCAGGG G UUGGGGAU	1969	ATCCCCAA GGCTAGCTACAACGA CCCTGAAA	4297
2926	GGUUGGG A UCUGUGCC	1970	GGCACAGA GGCTAGCTACAACGA CCCCACC	4298
2930	GGGAUCU G UGCCGGUG	1971	CACCGGCA GGCTAGCTACAACGA AGATCCCC	4299
2932	GGAUCUGU G CCGGUGGC	1972	GCCACCGG GGCTAGCTACAACGA ACAGATCC	4300
2936	CUGUGCCG G UGGCUCUG	1973	CAGAGCCA GGCTAGCTACAACGA CGGCACAG	4301
2939	UGCCGGUG G CUCUGGUC	1974	GACCAGAG GGCTAGCTACAACGA CACCGGCA	4302
2945	UGGCUCUG G UCUCUGCU	1975	AGCAGAGA GGCTAGCTACAACGA CAGAGCCA	4303
2951	UGGUCUCU G CUGGGAGC	1976	GCTCCAG GGCTAGCTACAACGA AGAGACCA	4304
2958	UGCUGGA G CCUUCUUG	1977	CAAGAAGG GGCTAGCTACAACGA TCCCAGCA	4305
2967	CCUUCUUG G CGGUGAGA	1978	TCTCACCG GGCTAGCTACAACGA CAAGAAGG	4306
2970	UCUUGGCG G UGAGAGGC	1979	GCCTCTCA GGCTAGCTACAACGA CGCCAAGA	4307
2977	GGUGAGAG G CAUACCU	1980	AGGTGATG GGCTAGCTACAACGA CTCTCACC	4308
2979	UGAGAGGC A UCACUUU	1981	AAAGGTGA GGCTAGCTACAACGA GCCTCTCA	4309
2982	GAGGAUC A CCUUCU	1982	AGGAAAGG GGCTAGCTACAACGA GATGCCTC	4310
2992	CUUUCUG A CUUGCUC	1983	GGAGCAAG GGCTAGCTACAACGA CAGGAAAG	4311
2996	CCUGACU G CUCCCAGC	1984	GCTGGGAG GGCTAGCTACAACGA AAGTCAGG	4312
3003	UGCUCCCA G CGUGAAAU	1985	ATTTACAG GGCTAGCTACAACGA TGGGAGCA	4313
3005	CUCCCAGC G UGAAAU	1986	GCATTTCA GGCTAGCTACAACGA GCTGGGAG	4314
3010	AGCGUGAA A UGCACCUG	1987	CAGGTGCA GGCTAGCTACAACGA TTCACGCT	4315
3012	CGUGAAAU G CACCUGCC	1988	GGCAGGTG GGCTAGCTACAACGA ATTTACAG	4316
3014	UGAAAU G CCUGCCAA	1989	TTGGCAGG GGCTAGCTACAACGA GCATTTCA	4317
3018	AUGCACCU G CCAAGAAU	1990	ATTCTTGG GGCTAGCTACAACGA AGGTGCAT	4318
3025	UGCCAAGA A UGGCAGAC	1991	GTCTGCCA GGCTAGCTACAACGA TCTTGGCA	4319
3028	CAAGAAUG G CAGACUA	1992	TATGTCG GGCTAGCTACAACGA CATTCTTG	4320
3032	AAUGGCAG A CAUAGGGA	1993	TCCCTATG GGCTAGCTACAACGA CTGCCATT	4321

3034	UGGCAGAC A UAGGGACC	1994	GGTCCCTA GGCTAGCTACAACGA GTCTGCCA	4322
3040	ACAUAGGG A CCCC GCCU	1995	AGGCGGGG GGCTAGCTACAACGA CCCTATGT	4323
3045	GGGACCCC G CCUCCUGG	1996	CCAGGAGG GGCTAGCTACAACGA GGGGTCCC	4324
3054	CCUCCUGG G CCUUCACA	1997	TGTGAAGG GGCTAGCTACAACGA CCAGGAGG	4325
3060	GGGCCUUC A CAUGCCCA	1998	TGGGCATG GGCTAGCTACAACGA GAAGGCCC	4326
3062	GCCUUCAC A UGCCCAGU	1999	ACTGGGCA GGCTAGCTACAACGA GTGAAGGC	4327
3064	CUUCACAU G CCCAGUUU	2000	AAACTGGG GGCTAGCTACAACGA ATGTGAAG	4328
3069	CAUGCCCA G UUUUCUUC	2001	GAAGAAAA GGCTAGCTACAACGA TGGGCATG	4329
3079	UUUCUUCG G CUCUGUGG	2002	CCACAGAG GGCTAGCTACAACGA CGAAGAAA	4330
3084	UCGGCUCU G UGGCCUGA	2003	TCAGGCCA GGCTAGCTACAACGA AGAGCCGA	4331
3087	GCUCUGUG G CCUGAAGC	2004	GCTTCAGG GGCTAGCTACAACGA CACAGAGC	4332
3094	GGCCUGAA G CGGUCUGU	2005	ACAGACCG GGCTAGCTACAACGA TTCAGGCC	4333
3097	CUGAAGCG G UCUGUGGA	2006	TCCACAGA GGCTAGCTACAACGA CGCTTCAG	4334
3101	AGCGGUCU G UGGACCUU	2007	AAGGTCCA GGCTAGCTACAACGA AGACCGCT	4335
3105	GUCUGUGG A CCUUGGAA	2008	TTCCAAGG GGCTAGCTACAACGA CCACAGAC	4336
3114	CCUUGGAA G UAGGGCUC	2009	GAGCCCTA GGCTAGCTACAACGA TTCCAAGG	4337
3119	GAAGUAGG G CUCCAGCA	2010	TGCTGGAG GGCTAGCTACAACGA CCTACTTC	4338
3125	GGGCUCCA G CACCGACU	2011	AGTCGGTG GGCTAGCTACAACGA TGGAGCCC	4339
3127	GCUCCAGC A CCGACUGG	2012	CCAGTCGG GGCTAGCTACAACGA GCTGGAGC	4340
3131	CAGCACCG A CUGGCCUC	2013	GAGGCCAG GGCTAGCTACAACGA CGGTGCTG	4341
3135	ACCGACUG G CCUCAGGC	2014	GCCTGAGG GGCTAGCTACAACGA CAGTCGGT	4342
3142	GGCCUCAG G CCUCUGCC	2015	GGCAGAGG GGCTAGCTACAACGA CTGAGGCC	4343
3148	AGGCCUCU G CCUCAUUG	2016	CAATGAGG GGCTAGCTACAACGA AGAGGCCT	4344
3153	UCUGCCUC A UUGGUGGU	2017	ACCACCAA GGCTAGCTACAACGA GAGGCAGA	4345
3157	CCUCAUUG G UGGUCGGG	2018	CCCGACCA GGCTAGCTACAACGA CAATGAGG	4346
3160	CAUUGGUG G UCGGGUAG	2019	CTACCCGA GGCTAGCTACAACGA CACCAATG	4347
3165	GUGGUCGG G UAGCGGCC	2020	GGCCGCTA GGCTAGCTACAACGA CCGACCAC	4348
3168	GUCGGGUA G CGGCCAGU	2021	ACTGGCCG GGCTAGCTACAACGA TACCCGAC	4349
3171	GGGUAGCG G CCAGUAGG	2022	CCTACTGG GGCTAGCTACAACGA CGCTACCC	4350
3175	AGCGGCCA G UAGGGCGU	2023	ACGCCCTA GGCTAGCTACAACGA TGGCCGCT	4351
3180	CCAGUAGG G CGUGGGAG	2024	CTCCACAG GGCTAGCTACAACGA CCTACTGG	4352
3182	AGUAGGGC G UGGGAGCC	2025	GGCTCCA GGCTAGCTACAACGA GCCCTACT	4353
3188	GCGUGGGA G CCUGGCCA	2026	TGGCCAGG GGCTAGCTACAACGA TCCCACGC	4354
3193	GGAGCCUG G CCAUCCCU	2027	AGGGATGG GGCTAGCTACAACGA CAGGCTCC	4355
3196	GCCUGGCC A UCCUGGCC	2028	GGCAGGGA GGCTAGCTACAACGA GGCCAGGC	4356
3202	CCAUCCCU G CCUCCUGG	2029	CCAGGAGG GGCTAGCTACAACGA AGGGATGG	4357
3212	CUCCUGGA G UGGACGAG	2030	CTCGTCCA GGCTAGCTACAACGA TCCAGGAG	4358
3216	UGGAGUGG A CGAGGUUG	2031	CAACCTCG GGCTAGCTACAACGA CCACTCCA	4359
3221	UGGACGAG G UUGGCAGC	2032	GCTGCCAA GGCTAGCTACAACGA CTCGTCCA	4360
3225	CGAGGUUG G CAGCUGGU	2033	ACCAGCTG GGCTAGCTACAACGA CAACCTCG	4361
3228	GGUUGGCA G CUGGUCCG	2034	CGGACCAG GGCTAGCTACAACGA TGCCAACC	4362
3232	GGCAGCUG G UCCGUCUG	2035	CAGACGGA GGCTAGCTACAACGA CAGCTGCC	4363
3236	GCUGGUCC G UCUGCUCC	2036	GGAGCAGA GGCTAGCTACAACGA GGACCAGC	4364
3240	GUCCGUCU G CUCCUGCC	2037	GGCAGGAG GGCTAGCTACAACGA AGACGGAC	4365
3246	CUGCUCCU G CCCCACUC	2038	GAGTGGGG GGCTAGCTACAACGA AGGAGCAG	4366
3251	CCUGCCCC A CUCUCCCC	2039	GGGGAGAG GGCTAGCTACAACGA GGGGCAGG	4367
3261	UCUCCCCC G CCCCUGCC	2040	GGCAGGGG GGCTAGCTACAACGA GGGGGAGA	4368
3267	CCGCCCCU G CCCUACCC	2041	GGTGAGGG GGCTAGCTACAACGA AGGGGCGG	4369
3273	CUGCCCU C A CCCUACCC	2042	GGGTAGGG GGCTAGCTACAACGA GAGGGCAG	4370
3278	CUCACCCU A CCCUUGCC	2043	GGCAAGGG GGCTAGCTACAACGA AGGGTGAG	4371
3284	CUACCCUU G CCCCACGC	2044	GCGTGGGG GGCTAGCTACAACGA AAGGGTAG	4372
3289	CUUGCCCC A CGCCUGCC	2045	GGCAGGCG GGCTAGCTACAACGA GGGGCAAG	4373

3291	UGCCCCAC G CCUGCCUC	2046	GAGGCAGG GGCTAGCTACAACGA GTGGGGCA	4374
3295	CCACGCCU G CCUCAUGG	2047	CCATGAGG GGCTAGCTACAACGA AGGCGTGG	4375
3300	CCUGCCUC A UGGCUGGU	2048	ACCAGCCA GGCTAGCTACAACGA GAGGCAGG	4376
3303	GCCUCAUG G CUGGUUGC	2049	GCAACCAG GGCTAGCTACAACGA CATGAGGC	4377
3307	CAUGGCUG G UUGCUCUU	2050	AAGAGCAA GGCTAGCTACAACGA CAGCCATG	4378
3310	GGCUGGUU G CUCUUGGA	2051	TCCAAGAG GGCTAGCTACAACGA AACCAGCC	4379
3319	CUCUUGGA G CCUGGUAG	2052	CTACCAGG GGCTAGCTACAACGA TCCAAGAG	4380
3324	GGAGCCUG G UAGUGUCA	2053	TGACACTA GGCTAGCTACAACGA CAGGCTCC	4381
3327	GCCUGGUA G UGUCACUG	2054	CAGTGACA GGCTAGCTACAACGA TACCAGGC	4382
3329	CUGGUAGU G UCACUGGC	2055	GCCAGTGA GGCTAGCTACAACGA ACTACCAG	4383
3332	GUAGUGUC A CUGGCUCA	2056	TGAGCCAG GGCTAGCTACAACGA GACACTAC	4384
3336	UGUCACUG G CUCAGCCU	2057	AGGTGAG GGCTAGCTACAACGA CAGTGACA	4385
3341	CUGGCUCA G CCUUGCUG	2058	CAGCAAGG GGCTAGCTACAACGA TGAGCCAG	4386
3346	UCAGCCUU G CUGGGUUAU	2059	ATACCCAG GGCTAGCTACAACGA AAGGCTGA	4387
3351	CUUGCUGG G UAUACACA	2060	TGTGTATA GGCTAGCTACAACGA CCAGCAAG	4388
3353	UGCUGGGU A UACACAGG	2061	CCTGTGTA GGCTAGCTACAACGA ACCCAGCA	4389
3355	CUGGGUUAU A CACAGGCU	2062	AGCCTGTG GGCTAGCTACAACGA ATACCCAG	4390
3357	GGGUUAUAC A CAGGCUCU	2063	AGAGCCTG GGCTAGCTACAACGA GTATACCC	4391
3361	AUACACAG G CUCUGCCA	2064	TGGCAGAG GGCTAGCTACAACGA CTGTGTAT	4392
3366	CAGGCUCU G CCACCCAC	2065	GTGGGTGG GGCTAGCTACAACGA AGAGCCTG	4393
3369	GCUCUGCC A CCCACUCU	2066	AGAGTGGG GGCTAGCTACAACGA GGCAGAGC	4394
3373	UGCCACCC A CUCUGCUC	2067	GAGCAGAG GGCTAGCTACAACGA GGGTGGCA	4395
3378	CCCACUCU G CUCCAAGG	2068	CCTTGGAG GGCTAGCTACAACGA AGAGTGGG	4396
3388	UCCAAGGG G CUUGCCCU	2069	AGGGCAAG GGCTAGCTACAACGA CCCTTGGGA	4397
3392	AGGGGCUU G CCCUGCCU	2070	AGGCAGGG GGCTAGCTACAACGA AAGCCCCT	4398
3397	CUUGCCCU G CCUUGGGC	2071	GCCCAAGG GGCTAGCTACAACGA AGGGCAAG	4399
3404	UGCCUUGG G CCAAGUUC	2072	GAACCTGG GGCTAGCTACAACGA CCAAGGCA	4400
3409	UGGGCCAA G UUCUAGGU	2073	ACCTAGAA GGCTAGCTACAACGA TTGGCCCA	4401
3416	AGUUCUAG G UCUGGCA	2074	TGGCCAGA GGCTAGCTACAACGA CTAGAACT	4402
3421	UAGGUCUG G CCACAGCC	2075	GGCTGTGG GGCTAGCTACAACGA CAGACCTA	4403
3424	GUCUGGCC A CAGCCACA	2076	TGTGGCTG GGCTAGCTACAACGA GGCCAGAC	4404
3427	UGGCCACA G CCACAGAC	2077	GTCTGTGG GGCTAGCTACAACGA TGTGGCCA	4405
3430	CCACAGCC A CAGACAGC	2078	GCTGTCTG GGCTAGCTACAACGA GGCTGTGG	4406
3434	AGCCACAG A CAGCUCAG	2079	CTGAGCTG GGCTAGCTACAACGA CTGTGGCT	4407
3437	CACAGACA G CUCAGUCC	2080	GGACTGAG GGCTAGCTACAACGA TGTCTGTG	4408
3442	ACAGCUCA G UCCCCUGU	2081	ACAGGGGA GGCTAGCTACAACGA TGAGCTGT	4409
3449	AGUCCCCU G UGUGUCA	2082	TGACCACA GGCTAGCTACAACGA AGGGGACT	4410
3451	UCCCCUGU G UGGUCAUC	2083	GATGACCA GGCTAGCTACAACGA ACAGGGGA	4411
3454	CCUGUGUG G UCAUCCUG	2084	CAGGATGA GGCTAGCTACAACGA CACACAGG	4412
3457	GUGUGGUC A UCCUGGCU	2085	AGCCAGGA GGCTAGCTACAACGA GACCACAC	4413
3463	UCAUCCUG G CUUCUGCU	2086	AGCAGAAG GGCTAGCTACAACGA CAGGATGA	4414
3469	UGGCUUCU G CUGGGGGC	2087	GCCCCCAG GGCTAGCTACAACGA AGAAGCCA	4415
3476	UGCUGGGG G CCCACAGC	2088	GCTGTGGG GGCTAGCTACAACGA CCCCAGCA	4416
3480	GGGGGCCC A CAGCGCCC	2089	GGGCGCTG GGCTAGCTACAACGA GGGCCCCC	4417
3483	GGCCACA G CGCCCUG	2090	CAGGGGCG GGCTAGCTACAACGA TGTGGGCC	4418
3485	CCCACAGC G CCCUGGU	2091	ACCAGGGG GGCTAGCTACAACGA GCTGTGGG	4419
3492	CGCCCUG G UGCCCUC	2092	GAGGGGCA GGCTAGCTACAACGA CAGGGGCG	4420
3494	CCCCUGGU G CCCUCCC	2093	GGGAGGGG GGCTAGCTACAACGA ACCAGGGG	4421
3511	CUCCCAGG G CCCGGGUU	2094	AACCCGGG GGCTAGCTACAACGA CCTGGGAG	4422
3517	GGGCCCCG G UUGAGGCU	2095	AGCCTCAA GGCTAGCTACAACGA CCGGGCCC	4423
3523	GGGUUGAG G CUGGGCCA	2096	TGGCCAG GGCTAGCTACAACGA CTCAACCC	4424
3528	GAGGCUGG G CCAGGCC	2097	GGGCCTGG GGCTAGCTACAACGA CCAGCCTC	4425

3533	UGGGCCAG G CCCUCUGG	2098	CCAGAGGG GGCTAGCTACAACGA CTGGCCCA	4426
3543	CCUCUGGG A CGGGGACU	2099	AGTCCCCG GGCTAGCTACAACGA CCCAGAGG	4427
3549	GGACGGGG A CUUGUGCC	2100	GGCACAAG GGCTAGCTACAACGA CCCCGTCC	4428
3553	GGGGACUU G UGCCCUGU	2101	ACAGGGCA GGCTAGCTACAACGA AAGTCCCC	4429
3555	GGACUUGU G CCCUGUCA	2102	TGACAGGG GGCTAGCTACAACGA ACAAGTCC	4430
3560	UGUGCCCU G UCAGGGUU	2103	AACCTTGA GGCTAGCTACAACGA AGGGCACA	4431
3566	CUGUCAGG G UUCCCUAU	2104	ATAGGGAA GGCTAGCTACAACGA CCTGACAG	4432
3573	GGUUCUU A UCCUGAG	2105	CTCAGGGA GGCTAGCTACAACGA AGGGAACC	4433
3582	UCCUGAG G UUGGGGA	2106	TCCCCAA GGCTAGCTACAACGA CTCAGGGA	4434
3593	GGGGAGA G CUAGCAGG	2107	CCTGCTAG GGCTAGCTACAACGA TCTCCCC	4435
3597	GAGAGCUA G CAGGGCAU	2108	ATGCCCTG GGCTAGCTACAACGA TAGCTCTC	4436
3602	CUAGCAGG G CAUGCCGC	2109	GCGGCATG GGCTAGCTACAACGA CCTGCTAG	4437
3604	AGCAGGGC A UGCCGUG	2110	CAGCGGCA GGCTAGCTACAACGA GCCCTGCT	4438
3606	CAGGGCAU G CCGCUGGC	2111	GCCAGCGG GGCTAGCTACAACGA ATGCCCTG	4439
3609	GGCAUGCC G CUGGCUGG	2112	CCAGCCAG GGCTAGCTACAACGA GGCATGCC	4440
3613	UGCCGUG G CUGGCCAG	2113	CTGGCCAG GGCTAGCTACAACGA CAGCGGCA	4441
3617	GCUGGCUG G CCAGGGCU	2114	AGCCCTGG GGCTAGCTACAACGA CAGCCAGC	4442
3623	UGGCCAGG G CUGCAGGG	2115	CCCTGCAG GGCTAGCTACAACGA CCTGGCCA	4443
3626	CCAGGGCU G CAGGGACA	2116	TGTCCCTG GGCTAGCTACAACGA AGCCCTGG	4444
3632	CUGCAGGG A CACUCCCC	2117	GGGGAGTG GGCTAGCTACAACGA CCCTGCAG	4445
3634	GCAGGGAC A CUCCCCU	2118	AGGGGGAG GGCTAGCTACAACGA GTCCCTGC	4446
3646	CCCCUUU G UCCAGGGA	2119	TCCCTGGA GGCTAGCTACAACGA AAAAGGGG	4447
3655	UCCAGGGA A UACCACAC	2120	GTGTGGTA GGCTAGCTACAACGA TCCCTGGA	4448
3657	CAGGGAAU A CCACACUC	2121	GAGTGTGG GGCTAGCTACAACGA ATTCCCTG	4449
3660	GGAAUACC A CACUCGCC	2122	GGCGAGTG GGCTAGCTACAACGA GGTATTCC	4450
3662	AAUACCAC A CUCGCCU	2123	AGGGCGAG GGCTAGCTACAACGA GTGGTATT	4451
3666	CCACACUC G CCCUUCUC	2124	GAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4452
3679	UCUCUCCA G CGAACACC	2125	GGTGTTCG GGCTAGCTACAACGA TGGAGAGA	4453
3683	UCCAGCGA A CACCACAC	2126	GTGTGGTG GGCTAGCTACAACGA TCGCTGGA	4454
3685	CAGCGAAC A CCACACUC	2127	GAGTGTGG GGCTAGCTACAACGA GTTCGCTG	4455
3688	CGAACACC A CACUCGCC	2128	GGCGAGTG GGCTAGCTACAACGA GGTGTTCG	4456
3690	AACACCAC A CUCGCCU	2129	AGGGCGAG GGCTAGCTACAACGA GTGGTGT	4457
3694	CCACACUC G CCCUUCUC	2124	GAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4452
3711	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3713	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCTTG	4459
3716	GGGACGCC A CACUCCCC	2132	GGGGAGTG GGCTAGCTACAACGA GGCGTCCC	4460
3718	GACGCCAC A CUCCCCU	2133	AGGGGGAG GGCTAGCTACAACGA GTGGCGTC	4461
3730	CCCCUUCU G UCCAGGGG	2134	CCCCTGGA GGCTAGCTACAACGA AGAAGGGG	4462
3739	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3741	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCTTG	4459
3744	GGGACGCC A CACUCCCC	2132	GGGGAGTG GGCTAGCTACAACGA GGCGTCCC	4460
3746	GACGCCAC A CUCCCCU	2133	AGGGGGAG GGCTAGCTACAACGA GTGGCGTC	4461
3767	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3769	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCTTG	4459
3772	GGGACGCC A CACUCGCC	2135	GGCGAGTG GGCTAGCTACAACGA GGCGTCCC	4463
3774	GACGCCAC A CUCGCCU	2136	AGGGCGAG GGCTAGCTACAACGA GTGGCGTC	4464
3778	CCACACUC G CCCUUCUC	2124	GAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4452
3795	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3797	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCTTG	4459
3800	GGGACGCC A CACUCGCC	2135	GGCGAGTG GGCTAGCTACAACGA GGCGTCCC	4463
3802	GACGCCAC A CUCGCCU	2136	AGGGCGAG GGCTAGCTACAACGA GTGGCGTC	4464
3806	CCACACUC G CCCUUCUC	2124	GAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4452

3823	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3825	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
3828	GGGACGCC A CACUCGCC	2135	GGCGAGTG GGCTAGCTACAACGA GGCGTCCC	4463
3830	GACGCCAC A CUCGCCCU	2136	AGGGCGAG GGCTAGCTACAACGA GTGGCGTC	4464
3834	CCACACUC G CCCUUCUG	2137	CAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4465
3842	GCCCUUCU G UCCAGGGG	2138	CCCCTGGA GGCTAGCTACAACGA AGAAGGGC	4466
3851	UCCAGGGG A CGCCACAC	2130	GTGTGGCG GGCTAGCTACAACGA CCCCTGGA	4458
3853	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
3856	GGGACGCC A CACUCGCC	2135	GGCGAGTG GGCTAGCTACAACGA GGCGTCCC	4463
3858	GACGCCAC A CUCGCCCU	2136	AGGGCGAG GGCTAGCTACAACGA GTGGCGTC	4464
3862	CCACACUC G CCCUUCUC	2124	GAGAAGGG GGCTAGCTACAACGA GAGTGTGG	4452
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3881	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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3912	GGGACGCC A CACUCCCC	2132	GGGGAGTG GGCTAGCTACAACGA GGCGTCCC	4460
3914	GACGCCAC A CUCGCCCU	2133	AGGGGGAG GGCTAGCTACAACGA GTGGCGTC	4461
3926	CCCUUCU G UCCAGGGG	2134	CCCCTGGA GGCTAGCTACAACGA AGAAGGGG	4462
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3937	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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3942	GACGCCAC A CUCGCCCU	2133	AGGGGGAG GGCTAGCTACAACGA GTGGCGTC	4461
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3993	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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4049	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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4105	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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4166	GACGCCAC A CUCCCCU	2133	AGGGGGAG GGCTAGCTACAACGA GTGGCGTC	4461
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4189	CAGGGGAC G CCACACUC	2131	GAGTGTGG GGCTAGCTACAACGA GTCCCCTG	4459
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4438	CCCCAGCA G CCUCCGAG	2149	CTCGGAGG GGCTAGCTACAACGA TGCTGGGG	4476
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4794	CGGGAACC A CUGGACAA	2226	TTGTCCAG GGCTAGCTACAACGA GGTTCCTG	4553
4799	ACCACUGG A CAACCUGG	2227	CCAGTTTG GGCTAGCTACAACGA CCAGTGGT	4554
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4904	CCCCUCAU G CCGCUAGG	2248	CCTAGCGG GGCTAGCTACAACGA ATGAGGGG	4575
4907	CUCAUGCC G CUAGGCCU	2249	AGGCCTAG GGCTAGCTACAACGA GGATGAG	4576
4912	GCCGCUAG G CCUUGGCC	2250	GGCCAAGG GGCTAGCTACAACGA CTAGCGGC	4577

4918	AGGCCUUG G CCUCGGGG	2251	CCCCGAGG GGCTAGCTACAACGA CAAGGCCT	4578
4927	CCUCGGGG A CAGCCAG	2252	CTGGGCTG GGCTAGCTACAACGA CCCGAGG	4579
4930	CGGGGACA G CCCAGCUA	2253	TAGCTGGG GGCTAGCTACAACGA TGTCCCCG	4580
4935	ACAGCCCA G CUAGGCCA	2254	TGGCCTAG GGCTAGCTACAACGA TGGGCTGT	4581
4940	CCAGCUAG G CCAGUGUG	2255	CACACTGG GGCTAGCTACAACGA CTAGCTGG	4582
4944	CUAGCCCA G UGUGUGGC	2256	GCCACACA GGCTAGCTACAACGA TGGCCTAG	4583
4946	AGGCCAGU G UGUGGCAG	2257	CTGCCACA GGCTAGCTACAACGA ACTGGCCT	4584
4948	GCCAGUGU G UGGCAGGA	2258	TCCTGCCA GGCTAGCTACAACGA ACACTGGC	4585
4951	AGUGUGUG G CAGGACCA	2259	TGGTCCTG GGCTAGCTACAACGA CACACACT	4586
4956	GUGGCAGG A CCAGGCC	2260	GGGCCTGG GGCTAGCTACAACGA CCTGCCAC	4587
4961	AGGACCAG G CCCCCAUG	2261	CATGGGGG GGCTAGCTACAACGA CTGGTCCT	4588
4967	AGGCCCC A UGUGGGAG	2262	CTCCACA GGCTAGCTACAACGA GGGGGCCT	4589
4969	CCCCCAU G UGGGAGCU	2263	AGCTCCA GGCTAGCTACAACGA ATGGGGGC	4590
4975	AUGUGGA G CUGACCC	2264	GGGGTCAG GGCTAGCTACAACGA TCCACAT	4591
4979	GGGAGCUG A CCCCUGG	2265	CCAAGGGG GGCTAGCTACAACGA CAGTCCC	4592
4989	CCCUUGG A UUCUGGAG	2266	CTCCAGAA GGCTAGCTACAACGA CCAAGGG	4593
4997	AUUCUGGA G CUGUGCUG	2267	CAGCACAG GGCTAGCTACAACGA TCCAGAAT	4594
5000	CUGGAGCU G UGCUGAUG	2268	CATCAGCA GGCTAGCTACAACGA AGTCCAG	4595
5002	GGAGCUGU G CUGAUGGG	2269	CCCATCAG GGCTAGCTACAACGA ACAGTCC	4596
5006	CUGUGCUG A UGGGCAGG	2270	CCTGCCCA GGCTAGCTACAACGA CAGCACAG	4597
5010	GCUGAUGG G CAGGGGAG	2271	CTCCCCTG GGCTAGCTACAACGA CCATCAGC	4598
5020	AGGGGAGA G CCAGCUCC	2272	GGAGCTGG GGCTAGCTACAACGA TCTCCCT	4599
5024	GAGAGCCA G CUCCUCCC	2273	GGGAGGAG GGCTAGCTACAACGA TGGCTCTC	4600
5044	GAGGGAGG G UCUUGAUG	2274	CATCAAGA GGCTAGCTACAACGA CCTCCCTC	4601
5050	GGGUCUUG A UGCCUGGG	2275	CCCAGGCA GGCTAGCTACAACGA CAAGACCC	4602
5052	GUCUUGAU G CCUGGGGU	2276	ACCCAGG GGCTAGCTACAACGA ATCAAGAC	4603
5059	UGCCUGGG G UUACCCGC	2277	GCGGGTAA GGCTAGCTACAACGA CCCAGGCA	4604
5062	CUGGGGUU A CCCGCAGA	2278	TCTGCGGG GGCTAGCTACAACGA AACCCAG	4605
5066	GGUUACCC G CAGAGGCC	2279	GGCCTCTG GGCTAGCTACAACGA GGGTAACC	4606
5072	CCGCAGAG G CCUGGGUG	2280	CACCCAGG GGCTAGCTACAACGA CTCTGCGG	4607
5078	AGGCCUGG G UGCCGGGA	2281	TCCCGGCA GGCTAGCTACAACGA CCAGGCCT	4608
5080	GCCUGGGU G CCGGGACG	2282	CGTCCCGG GGCTAGCTACAACGA ACCCAGGC	4609
5086	GUGCCGGG A CGCUCCCC	2283	GGGGAGCG GGCTAGCTACAACGA CCCGGCAC	4610
5088	GCCGGGAC G CUCCCCG	2284	CCGGGGAG GGCTAGCTACAACGA GTCCCGGC	4611
5096	GCUCCCCG G UUUGGCUG	2285	CAGCCAAA GGCTAGCTACAACGA CGGGGAGC	4612
5101	CCGGUUUG G CUGAAAGG	2286	CCTTTCAG GGCTAGCTACAACGA CAAACCGG	4613
5113	AAAGGAAA G CAGAUGUG	2287	CACATCTG GGCTAGCTACAACGA TTTCTTT	4614
5117	GAAAGCAG A UGUGGUCA	2288	TGACCACA GGCTAGCTACAACGA CTGCTTTC	4615
5119	AAGCAGAU G UGUUCAGC	2289	GCTGACCA GGCTAGCTACAACGA ATCTGCTT	4616
5122	CAGAUGUG G UCAGCUUC	2290	GAAGCTGA GGCTAGCTACAACGA CACATCTG	4617
5126	UGUGGUCA G CUUCUCCA	2291	TGGAGAAG GGCTAGCTACAACGA TGACCACA	4618
5134	GCUUCUCC A CUGAGCCC	2292	GGGCTCAG GGCTAGCTACAACGA GGAGAAGC	4619
5139	UCCACUGA G CCAUCUG	2293	CAGATGGG GGCTAGCTACAACGA TCAGTGGA	4620
5143	CUGAGCCC A UCUGGUCU	2294	AGACCAGA GGCTAGCTACAACGA GGGCTCAG	4621
5148	CCAUCUG G UCUUCCG	2295	CGGGAAGA GGCTAGCTACAACGA CAGATGGG	4622
5159	UUCCGGG G CUGGGCCC	2296	GGGCCAG GGCTAGCTACAACGA CCCGGGAA	4623
5164	GGGCGUGG G CCCCAG	2297	CTATGGGG GGCTAGCTACAACGA CCAGCCCC	4624
5169	UGGGCCCC A UAGAUCUG	2298	CAGATCTA GGCTAGCTACAACGA GGGGCCCC	4625
5173	CCCCAUG A UCUGGGUC	2299	GACCCAGA GGCTAGCTACAACGA CTATGGGG	4626
5179	AGAUCUGG G UCCUGUG	2300	CACAGGGA GGCTAGCTACAACGA CCAGATCT	4627
5185	GGGUCCCU G UGUGGCCC	2301	GGGCCACA GGCTAGCTACAACGA AGGGACCC	4628
5187	GUCCUCUGU G UGGCCCC	2302	GGGGGCCA GGCTAGCTACAACGA ACAGGGAC	4629

5190	CCUGUGUG G CCCCCUG	2303	CAGGGGGG GGCTAGCTACAACGA CACACAGG	4630
5199	CCCCCUG G UCUGAUGC	2304	GCATCAGA GGCTAGCTACAACGA CAGGGGGG	4631
5204	CUGGUCUG A UGCCGAGG	2305	CCTCGGCA GGCTAGCTACAACGA CAGACCAG	4632
5206	GGUCUGAU G CCGAGGAU	2306	ATCCTCGG GGCTAGCTACAACGA ATCAGACC	4633
5213	UGCCGAGG A UACCCUG	2307	CAGGGGTA GGCTAGCTACAACGA CCTCGGCA	4634
5215	CCGAGGAU A CCCCUGCA	2308	TGCAGGGG GGCTAGCTACAACGA ATCCTCGG	4635
5221	AUACCCCU G CAAACUGC	2309	GCAGTTTG GGCTAGCTACAACGA AGGGGTAT	4636
5225	CCCUGCAA A CUGCCAU	2310	ATTGGCAG GGCTAGCTACAACGA TTGCAGGG	4637
5228	UGCAAACU G CCAAUCCC	2311	GGGATTGG GGCTAGCTACAACGA AGTTTGCA	4638
5232	AACUGCCA A UCCCAGAG	2312	CTCTGGGA GGCTAGCTACAACGA TGGCAGTT	4639
5242	CCCAGAGG A CAAGACUG	2313	CAGTCTTG GGCTAGCTACAACGA CCTCTGGG	4640
5247	AGGACAAG A CUGGGAAG	2314	CTTCCCAG GGCTAGCTACAACGA CTTGTCTT	4641
5255	ACUGGGAA G UCCCUGCA	2315	TGCAGGGA GGCTAGCTACAACGA TTCCCAGT	4642
5261	AAGUCCCU G CAGGGAGA	2316	TCTCCCTG GGCTAGCTACAACGA AGGGACTT	4643
5270	CAGGGAGA G CCCAUCCC	2317	GGGATGGG GGCTAGCTACAACGA TCTCCCTG	4644
5274	GAGAGCCC A UCCCCGCA	2318	TGCGGGGA GGCTAGCTACAACGA GGGCTCTC	4645
5280	CCAUCCCC G CACCCUGA	2319	TCAGGGTG GGCTAGCTACAACGA GGGGATGG	4646
5282	AUCCCCGC A CCCUGACC	2320	GGTCAGGG GGCTAGCTACAACGA GCGGGGAT	4647
5288	GCACCCUG A CCCACAAG	2321	CTTGTGGG GGCTAGCTACAACGA CAGGGTGC	4648
5292	CCUGACCC A CAAGAGGG	2322	CCCTCTTG GGCTAGCTACAACGA GGGTCAGG	4649
5301	CAAGAGGG A CUCCUGCU	2323	AGCAGGAG GGCTAGCTACAACGA CCCTCTTG	4650
5307	GGACUCCU G CUGCCCAC	2324	GTGGGCAG GGCTAGCTACAACGA AGGAGTCC	4651
5310	CUCCUGCU G CCCACCAG	2325	CTGGTGGG GGCTAGCTACAACGA AGCAGGAG	4652
5314	UGCUGCCC A CCAGGCAU	2326	ATGCCTGG GGCTAGCTACAACGA GGGCAGCA	4653
5319	CCCACCAG G CAUCCUC	2327	GAGGGATG GGCTAGCTACAACGA CTGGTGGG	4654
5321	CACCAGGC A UCCCUCCA	2328	TGGAGGGA GGCTAGCTACAACGA GCCTGGTG	4655

Input Sequence = HUMRasH_mRNA. Cut Site = R/Y

Arm Length = 8. Core Sequence = GGCTAGCTACAACGA

HUMRasH_mRNA (Human c-Ha-ras1 proto-oncogene, spliced mRNA sequence; 5336 nt)

Table IV: Human HER2 DNzyme and Substrate Sequence

Pos	Substrate	Seq ID	DNzyme	Seq ID
9	AAGGGGAG G UAACCCUG	4656	CAGGGTTA GGCTAGCTACAACGA CTCCCCCTT	5644
12	GGGAGGUA A CCCUGGCC	4657	GGCCAGGG GGCTAGCTACAACGA TACCTCCC	5645
18	UAACCCUG G CCCCUUUG	4658	CAAAGGGG GGCTAGCTACAACGA CAGGGTTA	5646
27	CCCUUUG G UCGGGGCC	4659	GGCCCCGA GGCTAGCTACAACGA CAAAGGGG	5647
33	UGGUCGGG G CCCCGGGC	4660	GCCCCGGG GGCTAGCTACAACGA CCCGACCA	5648
40	GGCCCCGG G CAGCCGCG	4661	CGCGGCTG GGCTAGCTACAACGA CCGGGGCC	5649
43	CCCGGGCA G CCGCGCGC	4662	GCGCGCGG GGCTAGCTACAACGA TGCCCGGG	5650
46	GGGCAGCC G CGCGCCCC	4663	GGGGCGCG GGCTAGCTACAACGA GGCTGCCC	5651
48	GCAGCCGC G CGCCCCU	4664	AAGGGGCG GGCTAGCTACAACGA GCGGCTGC	5652
50	AGCCGCGC G CCCCUUCC	4665	GGAAGGGG GGCTAGCTACAACGA GCGCGGCT	5653
60	CCCUUCCC A CGGGGCC	4666	GGGCCCCG GGCTAGCTACAACGA GGAAGGGG	5654
65	CCCACGGG G CCCUUUAC	4667	GTAAAGGG GGCTAGCTACAACGA CCCGTGGG	5655
72	GGCCUUU A CUGCGCCG	4668	CGGCGCAG GGCTAGCTACAACGA AAAGGGCC	5656
75	CCUUUACU G CGCCGCGC	4669	GCGCGCGG GGCTAGCTACAACGA AGTAAAGG	5657
77	UUUACUGC G CCGCGCGC	4670	GCGCGCGG GGCTAGCTACAACGA GCAGTAAA	5658
80	ACUGCGCC G CGCGCCG	4671	CGGGCGCG GGCTAGCTACAACGA GCGCAGT	5659
82	UGCGCCGC G CGCCCGGC	4672	GCCGGGCG GGCTAGCTACAACGA GCGGCGCA	5660
84	CGCCGCGC G CCCGGCCC	4673	GGGCGGGG GGCTAGCTACAACGA GCGGCGCG	5661
89	CGCGCCCG G CCCCCACC	4674	GGTGGGGG GGCTAGCTACAACGA CGGGCGCG	5662
95	CGGCCCCC A CCCUUGC	4675	GCGAGGGG GGCTAGCTACAACGA GGGGGCCG	5663
102	CACCCUC G CAGCACC	4676	GGGTGCTG GGCTAGCTACAACGA GAGGGGTG	5664
105	CCCUCGCA G CACCCGCG	4677	GCGGGGTG GGCTAGCTACAACGA TGCGAGGG	5665
107	CUCGCAGC A CCCCAGCG	4678	GCGCGGGG GGCTAGCTACAACGA GCTGCGAG	5666
112	AGCACCCC G CGCCCGCG	4679	GCGGGGCG GGCTAGCTACAACGA GGGGTGCT	5667
114	CACCCGCG G CCCCAGCG	4680	GCGCGGGG GGCTAGCTACAACGA GCGGGGTG	5668
119	CGCGCCCC G CGCCUCC	4681	GGAGGGCG GGCTAGCTACAACGA GGGGCGCG	5669
121	CGCCCCGC G CCCUCCA	4682	TGGGAGGG GGCTAGCTACAACGA GCGGGGCG	5670
130	CCCUCCA G CCGGUCC	4683	GGACCCGG GGCTAGCTACAACGA TGGGAGGG	5671
135	CCAGCCGG G UCCAGCCG	4684	CGGCTGGA GGCTAGCTACAACGA CCGGCTGG	5672
140	CGGUCCA G CCGAGCC	4685	GGCTCCGG GGCTAGCTACAACGA TGGACCCG	5673
146	CAGCCGGA G CCAUGGGG	4686	CCCCATGG GGCTAGCTACAACGA TCCGGCTG	5674
149	CGGAGCC A UGGGGCCG	4687	CGGCCCA GGCTAGCTACAACGA GGCTCCGG	5675
154	GCCAUGGG G CCGAGCC	4688	GGCTCCGG GGCTAGCTACAACGA CCCATGGC	5676
160	GGGCCGGA G CCGCAGUG	4689	CACTGCGG GGCTAGCTACAACGA TCCGGCCC	5677
163	CCGAGCC G CAGUGAGC	4690	GCTCACTG GGCTAGCTACAACGA GGCTCCGG	5678
166	GAGCCGCA G UGAGCACC	4691	GGTGCTCA GGCTAGCTACAACGA TGCGGCTC	5679
170	CGCAGUGA G CACCAUGG	4692	CCATGGTG GGCTAGCTACAACGA TCACTGCG	5680
172	CAGUGAGC A CCAUGGAG	4693	CTCCATGG GGCTAGCTACAACGA GCTCACTG	5681
175	UGAGCACC A UGGAGCUG	4694	CAGCTCCA GGCTAGCTACAACGA GGTGCTCA	5682
180	ACCAUGGA G CUGGCGGC	4695	GCCGCCAG GGCTAGCTACAACGA TCCATGGT	5683
184	UGGAGCUG G CGGCCUUG	4696	CAAGGCCG GGCTAGCTACAACGA CAGCTCCA	5684
187	AGCUGGCG G CCUUGUGC	4697	GCACAAGG GGCTAGCTACAACGA CGCCAGCT	5685
192	GCGGCCUU G UGCGCUG	4698	CAGCGGCA GGCTAGCTACAACGA AAGGCCGC	5686
194	GGCCUUGU G CCGUGGG	4699	CCCAGCGG GGCTAGCTACAACGA ACAAGGCC	5687
197	CUUGUGCC G CUGGGGGC	4700	GCCCCCAG GGCTAGCTACAACGA GGCACAAG	5688
204	CGCUGGGG G CUCCUCCU	4701	AGGAGGAG GGCTAGCTACAACGA CCCAGCG	5689
214	UCCUCCUC G CCCUCUUG	4702	CAAGAGGG GGCTAGCTACAACGA GAGGAGGA	5690

222	GCCCUCUU G CCCCCCGG	4703	CCGGGGGG GGCTAGCTACAACGA AAGAGGGC	5691
232	CCCCCGGA G CCGCGAGC	4704	GCTCGCGG GGCTAGCTACAACGA TCCGGGGG	5692
235	CCGGAGCC G CGAGCACC	4705	GGTGCTCG GGCTAGCTACAACGA GGCTCCGG	5693
239	AGCCGCGA G CACCCAAG	4706	CTTGGGTG GGCTAGCTACAACGA TCGCCGGT	5694
241	CCGCGAGC A CCAAGUG	4707	CACTTGGG GGCTAGCTACAACGA GCTCGCGG	5695
247	GCACCCAA G UGUGCACC	4708	GGTGCA CA GGCTAGCTACAACGA TTGGGTGC	5696
249	ACCCAAGU G UGCACCGG	4709	CCGGTGCA GGCTAGCTACAACGA ACTTGGGT	5697
251	CCAAGUGU G CACCGGCA	4710	TGCCGGTG GGCTAGCTACAACGA ACACTTGG	5698
253	AAGUGUGC A CCGGCACA	4711	TGTGCCGG GGCTAGCTACAACGA GCACACTT	5699
257	GUGCACC G CACAGACA	4712	TGTCTGTG GGCTAGCTACAACGA CGGTGCAC	5700
259	GCACCGGC A CAGACAUG	4713	CATGTC TG GGCTAGCTACAACGA GCCGGTGC	5701
263	CGGCACAG A CAUGAAGC	4714	GCTTCATG GGCTAGCTACAACGA CTGTGCCG	5702
265	GCACAGAC A UGAAGCUG	4715	CAGCTTCA GGCTAGCTACAACGA GTCTGTGC	5703
270	GACAUGAA G CUGCGGCU	4716	AGCCGCAG GGCTAGCTACAACGA TTCATGTC	5704
273	AUGAAGCU G CGGCUCCC	4717	GGGAGCCG GGCTAGCTACAACGA AGCTTCAT	5705
276	AAGCUGCG G CUCCUGC	4718	GCAGGGAG GGCTAGCTACAACGA CGCAGCTT	5706
283	GGCUCCCU G CCAGUCCC	4719	GGGACTGG GGCTAGCTACAACGA AGGGAGCC	5707
287	CCUGCCA G UCCCGAGA	4720	TCTCGGGA GGCTAGCTACAACGA TGGCAGGG	5708
295	GUCCCGAG A CCCACCUG	4721	CAGGTGGG GGCTAGCTACAACGA CTCGGGAC	5709
299	CGAGACC A CCUGGACA	4722	TGTCCAGG GGCTAGCTACAACGA GGGTCTCG	5710
305	CCACCUGG A CAUGUCC	4723	GGAGCATG GGCTAGCTACAACGA CCAGGTGG	5711
307	ACCUGGAC A UGCUCCG	4724	GCGGAGCA GGCTAGCTACAACGA GTCCAGGT	5712
309	CUGGACAU G CUCCGCCA	4725	TGGCGGAG GGCTAGCTACAACGA ATGTCCAG	5713
314	CAUGUCC G CCACCUCU	4726	AGAGGTGG GGCTAGCTACAACGA GGAGCATG	5714
317	GCUCGCGC A CCUCUACC	4727	GGTAGAGG GGCTAGCTACAACGA GGCGGAGC	5715
323	CCACCUCU A CCAGGGCU	4728	AGCCCTGG GGCTAGCTACAACGA AGAGGTGG	5716
329	CUACCAGG G CUGCCAGG	4729	CCTGGCAG GGCTAGCTACAACGA CCTGGTAG	5717
332	CCAGGGCU G CCAGGUGG	4730	CCACCTGG GGCTAGCTACAACGA AGCCCTGG	5718
337	GCUGCCAG G UGUGCAG	4731	CTGCACCA GGCTAGCTACAACGA CTGGCAGC	5719
340	GCCAGGUG G UGCAGGGA	4732	TCCCTGCA GGCTAGCTACAACGA CACCTGGC	5720
342	CAGGUGGU G CAGGGAAA	4733	TTTCCCTG GGCTAGCTACAACGA ACCACCTG	5721
350	GCAGGGAA A CCUGGAAC	4734	GTTCCAGG GGCTAGCTACAACGA TTTCCCTG	5722
357	AACCUGGA A CUCACCUA	4735	TAGGTGAG GGCTAGCTACAACGA TCCAGGTT	5723
361	UGGAACUC A CCUACCUG	4736	CAGGTAGG GGCTAGCTACAACGA GAGTTCCA	5724
365	ACUCACCU A CCUGCCCA	4737	TGGGCAGG GGCTAGCTACAACGA AGGTGAGT	5725
369	ACCUACCU G CCCACCAA	4738	TTGGTGGG GGCTAGCTACAACGA AGGTAGGT	5726
373	ACCUGCCC A CCAAUGCC	4739	GGCATTGG GGCTAGCTACAACGA GGGCAGGT	5727
377	GCCCACCA A UGCCAGCC	4740	GGCTGGCA GGCTAGCTACAACGA TGGTGGGC	5728
379	CCACCAAU G CCAGCCUG	4741	CAGGCTGG GGCTAGCTACAACGA ATTGGTGG	5729
383	CAAUGCCA G CCUGUCCU	4742	AGGACAGG GGCTAGCTACAACGA TGGCATTG	5730
387	GCCAGCCU G UCCUCCU	4743	AGGAAGGA GGCTAGCTACAACGA AGGCTGGC	5731
396	UCCUCCU G CAGGAUUA	4744	ATATCCTG GGCTAGCTACAACGA AGGAAGGA	5732
401	CCUGCAGG A UAUCAGG	4745	CCTGGATA GGCTAGCTACAACGA CCTGCAGG	5733
403	UGCAGGAU A UCCAGGAG	4746	CTCCTGGA GGCTAGCTACAACGA ATCCTGCA	5734
412	UCCAGGAG G UGCAGGGC	4747	GCCCTGCA GGCTAGCTACAACGA CTCCTGGA	5735
414	CAGGAGGU G CAGGGCUA	4748	TAGCCCTG GGCTAGCTACAACGA ACCTCCTG	5736
419	GGUGCAGG G CUACGUGC	4749	GCACGTAG GGCTAGCTACAACGA CCTGCACC	5737
422	GCAGGGCU A CGUGCUCA	4750	TGAGCACG GGCTAGCTACAACGA AGCCCTGC	5738
424	AGGGCUAC G UGCUCAUC	4751	GATGAGCA GGCTAGCTACAACGA GTAGCCCT	5739
426	GGCUACGU G CUCAUCGC	4752	GCGATGAG GGCTAGCTACAACGA ACGTAGCC	5740
430	ACGUGCUC A UCGCUCAC	4753	GTGAGCGA GGCTAGCTACAACGA GAGCACGT	5741
433	UGCUCauc G CUCACAAC	4754	GTTGTGAG GGCTAGCTACAACGA GATGAGCA	5742

437	CAUCGCUC A CAACCAAG	4755	CTTGGTTG GGCTAGCTACAACGA GAGCGATG	5743
440	CGCUCACA A CCAAGUGA	4756	TCACTTGG GGCTAGCTACAACGA TGTGAGCG	5744
445	ACAACCAA G UGAGGCAG	4757	CTGCCTCA GGCTAGCTACAACGA TTGGTTGT	5745
450	CAAGUGAG G CAGGUCCC	4758	GGGACCTG GGCTAGCTACAACGA CTCACTTG	5746
454	UGAGGCAG G UCCCACUG	4759	CAGTGGGA GGCTAGCTACAACGA CTGCCTCA	5747
459	CAGGUCCC A CUGCAGAG	4760	CTCTGCAG GGCTAGCTACAACGA GGGACCTG	5748
462	GUCCCACU G CAGAGGCU	4761	AGCCTCTG GGCTAGCTACAACGA AGTGGGAC	5749
468	CUGCAGAG G CUGCGGAU	4762	ATCCGCAG GGCTAGCTACAACGA CTCTGCAG	5750
471	CAGAGGCU G CGGAUUGU	4763	ACAATCCG GGCTAGCTACAACGA AGCCTCTG	5751
475	GGCUGCGG A UUGUGCGA	4764	TCGCACAA GGCTAGCTACAACGA CCGCAGCC	5752
478	UGCGGAUU G UGCGAGGC	4765	GCCTCGCA GGCTAGCTACAACGA AATCCGCA	5753
480	CGGAUUGU G CGAGGCAC	4766	GTGCCTCG GGCTAGCTACAACGA ACAATCCG	5754
485	UGUGCGAG G CACCCAGC	4767	GCTGGGTG GGCTAGCTACAACGA CTCGCACA	5755
487	UGCGAGGC A CCCAGCUC	4768	GAGCTGGG GGCTAGCTACAACGA GCCTCGCA	5756
492	GGCACCCA G CUCUUUGA	4769	TCAAAGAG GGCTAGCTACAACGA TGGGTGCC	5757
503	CUUUGAGG A CAACUAUG	4770	CATAGTTG GGCTAGCTACAACGA CCTCAAAG	5758
506	UGAGGACA A CUAUGCCC	4771	GGGCATAG GGCTAGCTACAACGA TGTCTCA	5759
509	GGACAACU A UGCCCUGG	4772	CCAGGGCA GGCTAGCTACAACGA AGTTGTCC	5760
511	ACAACUUAU G CCCUGGCC	4773	GGCCAGGG GGCTAGCTACAACGA ATAGTTGT	5761
517	AUGCCUG G CCGUGCUA	4774	TAGCACGG GGCTAGCTACAACGA CAGGGCAT	5762
520	CCCUGGCC G UGCUAGAC	4775	GTCTAGCA GGCTAGCTACAACGA GGCCAGGG	5763
522	CUGGCCGU G CUAGACAA	4776	TTGTCTAG GGCTAGCTACAACGA ACGGCCAG	5764
527	CGUGCUAG A CAAUGGAG	4777	CTCCATTG GGCTAGCTACAACGA CTAGCACG	5765
530	GCUAGACA A UGGAGACC	4778	GGTCTCCA GGCTAGCTACAACGA TGTCTAGC	5766
536	CAAUGGAG A CCCGUGA	4779	TCAGCGGG GGCTAGCTACAACGA CTCCATTG	5767
540	GGAGACCC G CUGAACAA	4780	TTGTTTCA GGCTAGCTACAACGA GGGTCTCC	5768
545	CCCGCUGA A CAAUACCA	4781	TGGTATTG GGCTAGCTACAACGA TCAGCGGG	5769
548	GCUGAACA A UACCACCC	4782	GGGTGGTA GGCTAGCTACAACGA TGTTCAGC	5770
550	UGAACAAU A CCACCCCU	4783	AGGGGTGG GGCTAGCTACAACGA ATTGTTCA	5771
553	ACAAUACC A CCCUGUC	4784	GACAGGGG GGCTAGCTACAACGA GGTATTGT	5772
559	CCACCCCU G UCACAGGG	4785	CCCTGTGA GGCTAGCTACAACGA AGGGGTGG	5773
562	CCCUGUC A CAGGGGCC	4786	GGCCCTG GGCTAGCTACAACGA GACAGGGG	5774
568	UCACAGGG G CCUCCCCA	4787	TGGGGAGG GGCTAGCTACAACGA CCCTGTGA	5775
581	CCCAGGAG G CCUGCGGG	4788	CCCGCAGG GGCTAGCTACAACGA CTCCTGGG	5776
585	GGAGGCCU G CGGGAGCU	4789	AGCTCCCG GGCTAGCTACAACGA AGGCCTCC	5777
591	CUGCGGGA G CUGCAGCU	4790	AGCTGCAG GGCTAGCTACAACGA TCCGCGAG	5778
594	CGGGAGCU G CAGCUUCG	4791	CGAAGCTG GGCTAGCTACAACGA AGCTCCCG	5779
597	GAGCUGCA G CUUCGAAG	4792	CTTCGAAG GGCTAGCTACAACGA TGCAGCTC	5780
605	GCUUCGAA G CCUCACAG	4793	CTGTGAGG GGCTAGCTACAACGA TTCGAAGC	5781
610	GAAGCCUC A CAGAGAUC	4794	GATCTCTG GGCTAGCTACAACGA GAGGCTTC	5782
616	UCACAGAG A UCUUGAAA	4795	TTTCAAGA GGCTAGCTACAACGA CTCTGTGA	5783
631	AAGGAGGG G UCUUGAUC	4796	GATCAAGA GGCTAGCTACAACGA CCCTCCTT	5784
637	GGGUCUUG A UCCAGCGG	4797	CCGCTGGA GGCTAGCTACAACGA CAAGACCC	5785
642	UUGAUCCA G CGGAACCC	4798	GGGTTCOG GGCTAGCTACAACGA TGGATCAA	5786
647	CCAGCGGA A CCCCCAGC	4799	GCTGGGGG GGCTAGCTACAACGA TCCGCTGG	5787
654	AACCCCCA G CUCUGCUA	4800	TAGCAGAG GGCTAGCTACAACGA TGGGGGTT	5788
659	CCAGCUCU G CUACCAGG	4801	CCTGGTAG GGCTAGCTACAACGA AGAGCTGG	5789
662	GCUCUGCU A CCAGGACA	4802	TGTCCTGG GGCTAGCTACAACGA AGCAGAGC	5790
668	CUACCAGG A CACGAUUU	4803	AAATCGTG GGCTAGCTACAACGA CCTGGTAG	5791
670	ACCAGGAC A CGAUUUUG	4804	CAAAATCG GGCTAGCTACAACGA GTCCTGGT	5792
673	AGGACACG A UUUUGUGG	4805	CCACAAA GGCTAGCTACAACGA CGTGTCTT	5793
678	ACGAUUUU G UGGAAGGA	4806	TCCTTCCA GGCTAGCTACAACGA AAAATCGT	5794

686	GUGGAAGG A CAUCUCC	4807	GGAAGATG GGCTAGCTACAACGA CCTTCCAC	5795
688	GGAAGGAC A UCUUCCAC	4808	GTGGAAGA GGCTAGCTACAACGA GTCCTTCC	5796
695	CAUCUCC A CAAGAACA	4809	TGTTCTTG GGCTAGCTACAACGA GGAAGATG	5797
701	CCACAAGA A CAACCAGC	4810	GCTGGTTG GGCTAGCTACAACGA TCTTGTGG	5798
704	CAAGAACA A CCAGCUGG	4811	CCAGCTGG GGCTAGCTACAACGA TGTTCTTG	5799
708	AACAACCA G CUGGCUCU	4812	AGAGCCAG GGCTAGCTACAACGA TGGTTGTT	5800
712	ACCAGCUG G CUCUCACA	4813	TGTGAGAG GGCTAGCTACAACGA CAGCTGGT	5801
718	UGGCUCUC A CACUGAUA	4814	TATCAGTG GGCTAGCTACAACGA GAGAGCCA	5802
720	GCUCUCAC A CUGAUAGA	4815	TCTATCAG GGCTAGCTACAACGA GTGAGAGC	5803
724	UCACACUG A UAGACACC	4816	GGTGTCTA GGCTAGCTACAACGA CAGTGTGA	5804
728	ACUGAUAG A CACCAACC	4817	GGTTGGTG GGCTAGCTACAACGA CTATCAGT	5805
730	UGAUAGAC A CCAACCGC	4818	GCGGTTGG GGCTAGCTACAACGA GTCTATCA	5806
734	AGACACCA A CCGCUCUC	4819	GAGAGCGG GGCTAGCTACAACGA TGGTGTCT	5807
737	CACCAACC G CUCUCGGG	4820	CCCAGAGG GGCTAGCTACAACGA GGTGGTG	5808
745	GCUCUCGG G CCUGCCAC	4821	GTGGCAGG GGCTAGCTACAACGA CCGAGAGC	5809
749	UCGGGCCU G CCACCCCU	4822	AGGGGTGG GGCTAGCTACAACGA AGGCCCGA	5810
752	GGCCUGCC A CCCUGUU	4823	AACAGGGG GGCTAGCTACAACGA GGCAGGCC	5811
758	CCACCCCU G UUCUCCGA	4824	TCGGAGAA GGCTAGCTACAACGA AGGGGTGG	5812
766	GUUCUCCG A UGUGUAAG	4825	CTTACACA GGCTAGCTACAACGA CGGAGAAC	5813
768	UCUCCGAU G UGUAAGGG	4826	CCCTTACA GGCTAGCTACAACGA ATCGGAGA	5814
770	UCCGAUGU G UAAGGGCU	4827	AGCCCTTA GGCTAGCTACAACGA ACATCGGA	5815
776	GUGUAAGG G CUCCCGCU	4828	AGCGGGAG GGCTAGCTACAACGA CCTTACAC	5816
782	GGGCUCCC G CUGCUGGG	4829	CCCAGCAG GGCTAGCTACAACGA GGGAGCCC	5817
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797	GGGAGAGA G UUCUGAGG	4831	CCTCAGAA GGCTAGCTACAACGA TCTCTCCC	5819
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815	UUGUCAGA G CCUGACGC	4834	GCGTCAGG GGCTAGCTACAACGA TCTGACAA	5822
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824	CCUGACGC G CACUGUCU	4837	AGACAGTG GGCTAGCTACAACGA GCCTCAGG	5825
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839	CUGUGCCG G UGGCUGUG	4842	CACAGCCA GGCTAGCTACAACGA CGGCACAG	5830
842	UGCCGGUG G CUGUGCCC	4843	GGGCACAG GGCTAGCTACAACGA CACCGGCA	5831
845	CGGUGGCU G UGCCCUCU	4844	AGCGGGCA GGCTAGCTACAACGA AGCCACCG	5832
847	GUGGCUGU G CCCGUCG	4845	GCAGCGGG GGCTAGCTACAACGA ACAGCCAC	5833
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867	GGGCCACU G CCCACUGA	4850	TCAGTGGG GGCTAGCTACAACGA AGTGGCCC	5838
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875	GCCCACUG A CUGCUGCC	4852	GGCAGCAG GGCTAGCTACAACGA CAGTGGGC	5840
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891	CAUGAGCA G UGUGCUGC	4857	GCAGCACA GGCTAGCTACAACGA TGCTCATG	5845
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918	GGCCCCAA G CACUCUGA	4865	TCAGAGTG GGCTAGCTACAACGA TTGGGGCC	5853
920	CCCCAAGC A CUCUGACU	4866	AGTCAGAG GGCTAGCTACAACGA GCTTGGGG	5854
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956	CAACCACA G UGGCAUCU	4874	AGATGCCA GGCTAGCTACAACGA TGTGGTTG	5862
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1058	CGCCAGCU G UGUGACUG	4903	CAGTCACA GGCTAGCTACAACGA AGCTGGCG	5891
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1586	CCACCAUA A CACCCACC	5026	GGTGGGTG GGCTAGCTACAACGA TATGGTGG	6014
1588	ACCAUAAC A CCCACCUC	5027	GAGGTGGG GGCTAGCTACAACGA GTTATGGT	6015
1592	UAACACCC A CCUCUGCU	5028	AGCAGAGG GGCTAGCTACAACGA GGGTGTTA	6016
1598	CCACCUCU G CUUCGUGC	5029	GCACGAAG GGCTAGCTACAACGA AGAGGTGG	6017
1603	UCUGCUUC G UGCACACG	5030	CGTGTGCA GGCTAGCTACAACGA GAAGCAGA	6018
1605	UGCUCUGU G CACACGGU	5031	ACCGTGTG GGCTAGCTACAACGA ACGAAGCA	6019
1607	CUUCGUGC A CACGGUGC	5032	GCACCGTG GGCTAGCTACAACGA GCACGAAG	6020
1609	UCGUGCAC A CGUGCCCC	5033	GGGCACCG GGCTAGCTACAACGA GTGCACGA	6021
1612	UGCACACG G UGCCCUGG	5034	CCAGGGCA GGCTAGCTACAACGA CGTGTGCA	6022
1614	CACACGGU G CCCUGGGA	5035	TCCCAGGG GGCTAGCTACAACGA ACCGTGTG	6023
1622	GCCCUGGG A CCAGCUCU	5036	AGAGCTGG GGCTAGCTACAACGA CCCAGGGC	6024
1626	UGGGACCA G CUCUUUCG	5037	CGAAAGAG GGCTAGCTACAACGA TGGTCCCA	6025
1637	CUUUCGGA A CCCGCACC	5038	GGTGCGGG GGCTAGCTACAACGA TCCGAAAG	6026
1641	CGGAACCC G CACCAAGC	5039	GCTTGGTG GGCTAGCTACAACGA GGGTTCCG	6027
1643	GAACCCGC A CCAAGCUC	5040	GAGCTTGG GGCTAGCTACAACGA GCGGGTTC	6028
1648	CGCACCAA G CUCUGCUC	5041	GAGCAGAG GGCTAGCTACAACGA TTGGTGCG	6029
1653	CAAGCUCU G CUCCACAC	5042	GTGTGGAG GGCTAGCTACAACGA AGAGCTTG	6030
1658	UCUGCUCC A CACUGCCA	5043	TGGCAGTG GGCTAGCTACAACGA GGAGCAGA	6031
1660	UGCUCAC A CUGCCAAC	5044	GTTGGCAG GGCTAGCTACAACGA GTGGAGCA	6032
1663	UCCACACU G CCAACCGG	5045	CCGTTGGG GGCTAGCTACAACGA AGTGTGGA	6033
1667	CACUGCCA A CCGGCCAG	5046	CTGGCCGG GGCTAGCTACAACGA TGGCAGTG	6034
1671	GCCAACCG G CCAGAGGA	5047	TCCTCTGG GGCTAGCTACAACGA CGGTTGGC	6035
1679	GCCAGAGG A CGAGUGUG	5048	CACACTCG GGCTAGCTACAACGA CCTCTGGC	6036
1683	GAGGACGA G UGUGUGGG	5049	CCCACACA GGCTAGCTACAACGA TCGTCCTC	6037
1685	GGACGAGU G UGUGGGCG	5050	CGCCCACA GGCTAGCTACAACGA ACTCGTCC	6038
1687	ACGAGUGU G UGGGCGAG	5051	CTCGCCCA GGCTAGCTACAACGA AACTCGT	6039
1691	GUGUGUGG G CGAGGGCC	5052	GGCCCTCG GGCTAGCTACAACGA CCACACAC	6040
1697	GGGCGAGG G CCUGGCCU	5053	AGGCCAGG GGCTAGCTACAACGA CCTCGCCC	6041
1702	AGGGCCUG G CCUGCCAC	5054	GTGGCAGG GGCTAGCTACAACGA CAGGCCCT	6042
1706	CCUGGCCU G CCACCAGC	5055	GCTGGTGG GGCTAGCTACAACGA AGGCCAGG	6043
1709	GGCCUGCC A CCAGCUGU	5056	ACAGCTGG GGCTAGCTACAACGA GGCAGGCC	6044
1713	UGCCACCA G CUGUGCGC	5057	GCGCACAG GGCTAGCTACAACGA TGGTGGCA	6045
1716	CACCAGCU G UGCGCCCG	5058	CGGGCGCA GGCTAGCTACAACGA AGCTGGTG	6046
1718	CCAGCUGU G CGCCCAG	5059	CTCGGGCG GGCTAGCTACAACGA ACAGCTGG	6047
1720	AGCUGUGC G CCCGAGGG	5060	CCCTCGGG GGCTAGCTACAACGA GCACAGCT	6048
1728	GCCCAGGG G CACUGCUG	5061	CAGCAGTG GGCTAGCTACAACGA CCTCGGGC	6049
1730	CCGAGGGC A CUGCUGGG	5062	CCCAGCAG GGCTAGCTACAACGA GCCCTCGG	6050
1733	AGGGCACU G CUGGGGUC	5063	GACCCACG GGCTAGCTACAACGA AGTGCCCT	6051
1739	CUGCUGGG G UCCAGGGC	5064	GCCCTGGA GGCTAGCTACAACGA CCCAGCAG	6052
1746	GGUCCAGG G CCCACCCA	5065	TGGGTGGG GGCTAGCTACAACGA CCTGGACC	6053
1750	CAGGGCCC A CCCAGUGU	5066	ACACTGGG GGCTAGCTACAACGA GGGCCCTG	6054

1755	CCCACCCA G UGUGUCAA	5067	TTGACACA GGCTAGCTACAACGA TGGGTGGG	6055
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1759	CCCAGUGU G UCAACUGC	5069	GCAGTTGA GGCTAGCTACAACGA AACTGGG	6057
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1766	UGUCAACU G CAGCCAGU	5071	ACTGGCTG GGCTAGCTACAACGA AGTTGACA	6059
1769	CAACUGCA G CCAGUUC	5072	GGAACTGG GGCTAGCTACAACGA TGCAGTTG	6060
1773	UGCAGCCA G UUCUUCG	5073	CGAAGGAA GGCTAGCTACAACGA TGGCTGCA	6061
1784	CCUUCGGG G CCAGGAGU	5074	ACTCCTGG GGCTAGCTACAACGA CCGAAGG	6062
1791	GGCCAGGA G UGCGUGGA	5075	TCCACGCA GGCTAGCTACAACGA TCCTGGCC	6063
1793	CCAGGAGU G CGUGGAGG	5076	CCTCCACG GGCTAGCTACAACGA ACTCCTGG	6064
1795	AGGAGUGC G UGGAGGAA	5077	TTCTCCA GGCTAGCTACAACGA GCACTCCT	6065
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1805	GGAGGAAU G CCGAGUAC	5079	GTACTCGG GGCTAGCTACAACGA ATTCTCC	6067
1810	AAUGCCGA G UACUGCAG	5080	CTGCAGTA GGCTAGCTACAACGA TCGGCATT	6068
1812	UGCCGAGU A CUGCAGGG	5081	CCCTGCAG GGCTAGCTACAACGA ACTCGGCA	6069
1815	CGAGUACU G CAGGGGCU	5082	AGCCCCTG GGCTAGCTACAACGA AGTACTCG	6070
1821	CUGCAGGG G CUCCCCAG	5083	CTGGGGAG GGCTAGCTACAACGA CCCTGCAG	6071
1833	CCCAGGGA G UAUGUGAA	5084	TTACATA GGCTAGCTACAACGA TCCTGGG	6072
1835	CAGGGAGU A UGUGAAUG	5085	CATTCA GGCTAGCTACAACGA ACTCCCTG	6073
1837	GGGAGUAAU G UGAAUGCC	5086	GGCATTCA GGCTAGCTACAACGA ATACTCCC	6074
1841	GUAUGUGA A UGCCAGGC	5087	GCCTGGCA GGCTAGCTACAACGA TCACATAC	6075
1843	AUGUGAAU G CCAGGCAC	5088	GTGCCTGG GGCTAGCTACAACGA ATTACAT	6076
1848	AAUGCCAG G CACUGUUU	5089	AAACAGTG GGCTAGCTACAACGA CTGGCATT	6077
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1862	UUUGCCGU G CCACCCUG	5094	CAGGGTGG GGCTAGCTACAACGA ACGGCAAA	6082
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1872	CACCCUGA G UGUCAGCC	5096	GGCTGACA GGCTAGCTACAACGA TCAGGGTG	6084
1874	CCCUGAGU G UCAGCCCC	5097	GGGGCTGA GGCTAGCTACAACGA ACTCAGGG	6085
1878	GAGUGUCA G CCCCAGAA	5098	TTCTGGGG GGCTAGCTACAACGA TGACACTC	6086
1886	GCCCCAGA A UGGCUCAG	5099	CTGAGCCA GGCTAGCTACAACGA TCTGGGGC	6087
1889	CCAGAAUG G CUCAGUGA	5100	TCACTGAG GGCTAGCTACAACGA CATTCTGG	6088
1894	AUGGCUCA G UGACCUGU	5101	ACAGGTCA GGCTAGCTACAACGA TGAGCCAT	6089
1897	GCUCAGUG A CCUGUUUU	5102	AAAACAGG GGCTAGCTACAACGA CACTGAGC	6090
1901	AGUGACCU G UUUUGGAC	5103	GTCCAAA GGCTAGCTACAACGA AGGTCACT	6091
1908	UGUUUUGG A CCGGAGGC	5104	GCCTCCGG GGCTAGCTACAACGA CCAAACA	6092
1915	GACCGGAG G CUGACCAG	5105	CTGGTCAG GGCTAGCTACAACGA CTCCGGTC	6093
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1923	GCUGACCA G UGUGUGGC	5107	GCCACACA GGCTAGCTACAACGA TGGTCAGC	6095
1925	UGACCAGU G UGUGGCCU	5108	AGGCCACA GGCTAGCTACAACGA ACTGGTCA	6096
1927	ACCAGUGU G UGGCCUGU	5109	ACAGGCCA GGCTAGCTACAACGA AACTGGT	6097
1930	AGUGUGUG G CCUGUGCC	5110	GGCACAGG GGCTAGCTACAACGA CACACACT	6098
1934	UGUGGCCU G UGCCACU	5111	AGTGGGCA GGCTAGCTACAACGA AGGCCACA	6099
1936	UGGCCUGU G CCCACUAAU	5112	ATAGTGGG GGCTAGCTACAACGA ACAGGCCA	6100
1940	CUGUGCCC A CUAAAGG	5113	CCTTATAG GGCTAGCTACAACGA GGGCACAG	6101
1943	UGCCACU A UAAGGACC	5114	GGTCCTTA GGCTAGCTACAACGA AGTGGGCA	6102
1949	CUAAAGG A CCCUCCU	5115	AGGGAGGG GGCTAGCTACAACGA CCTTATAG	6103
1961	UCCUUCU G CGUGGCC	5116	GGGCCACG GGCTAGCTACAACGA AGAAGGGA	6104
1963	CCUUCUGC G UGGCCCGC	5117	GCGGGCCA GGCTAGCTACAACGA GCAGAAGG	6105
1966	UCUGCGUG G CCGCUGC	5118	GCAGCGGG GGCTAGCTACAACGA CACGCAGA	6106

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1973	GGCCCGCU G CCCAGCG	5120	CGCTGGGG GGCTAGCTACAACGA AGCGGGCC	6108
1979	CUGCCCCA G CGGUGUGA	5121	TCACACCG GGCTAGCTACAACGA TGGGGCAG	6109
1982	CCCAGCG G UGUGAAAC	5122	GTTTCACA GGCTAGCTACAACGA CGCTGGGG	6110
1984	CCAGCGGU G UGAAACCU	5123	AGGTTTCA GGCTAGCTACAACGA ACCGCTGG	6111
1989	GGUGUGAA A CCUGACCU	5124	AGGTCAGG GGCTAGCTACAACGA TTCACACC	6112
1994	GAAACCU G A CCUCUCCU	5125	AGGAGAGG GGCTAGCTACAACGA CAGGTTTC	6113
2003	CCUCUCCU A CAUGCCCA	5126	TGGGCATG GGCTAGCTACAACGA AGGAGAGG	6114
2005	UCUCCUAC A UGCCCAC	5127	GATGGGCA GGCTAGCTACAACGA GTAGGAGA	6115
2007	UCCUACAU G CCCAUCUG	5128	CAGATGGG GGCTAGCTACAACGA ATGTAGGA	6116
2011	ACAUGCCC A UCUGGAAG	5129	CTTCCAGA GGCTAGCTACAACGA GGGCATGT	6117
2019	AUCUGGAA G UUUCGAGA	5130	TCTGAAA GGCTAGCTACAACGA TTCCAGAT	6118
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2042	GGGCGCAU G CCAGCCUU	5135	AAGGCTGG GGCTAGCTACAACGA ATGCGCCC	6123
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2060	CCCCAUCA A CUGCACCC	5139	GGGTGCAG GGCTAGCTACAACGA TGATGGGG	6127
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2096	UGACAAGG G CUGCCCCG	5148	CGGGGCAG GGCTAGCTACAACGA CCTTGTCA	6136
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2226	AUCCGGAA G UACACGAU	5176	ATCGTGTA GGCTAGCTACAACGA TTCCGGAT	6164
2228	CCGGAAGU A CACGAUGC	5177	GCATCGTG GGCTAGCTACAACGA ACTTCCGG	6165
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2518	CCCCAUAU G UCUCCCGC	5243	GCGGGAGA GGCTAGCTACAACGA ATATGGGG	6231
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2534	CCUUCUGG G CAUCUGCC	5245	GGCAGATG GGCTAGCTACAACGA CCAGAAGG	6233
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2551	UGACAUC A CGGUGCAG	5250	CTGCACCG GGCTAGCTACAACGA GGATGTCA	6238
2554	CAUCCACG G UGCAGCUG	5251	CAGCTGCA GGCTAGCTACAACGA CGTGGATG	6239
2556	UCCACGGU G CAGCUGGU	5252	ACCAGCTG GGCTAGCTACAACGA ACCGTGGA	6240
2559	ACGGUGCA G CUGGUGAC	5253	GTCACCAG GGCTAGCTACAACGA TGCACCGT	6241
2563	UGCAGCUG G UGACACAG	5254	CTGTGTCA GGCTAGCTACAACGA CAGCTGCA	6242
2566	AGCUGGUG A CACAGCUU	5255	AAGCTGTG GGCTAGCTACAACGA CACCAGCT	6243
2568	CUGGUGAC A CAGCUUUA	5256	ATAAGCTG GGCTAGCTACAACGA GTCACCAG	6244
2571	GUGACACA G CUUAUGCC	5257	GGCATAAG GGCTAGCTACAACGA TGTGTAC	6245
2575	CACAGCUU A UGCCUUAU	5258	ATAGGGCA GGCTAGCTACAACGA AAGCTGTG	6246
2577	CAGCUUAU G CCCUAUGG	5259	CCATAGGG GGCTAGCTACAACGA ATAAGCTG	6247
2582	UAUGCCCU A UGGCUGCC	5260	GGCAGCCA GGCTAGCTACAACGA AGGGCATA	6248
2585	GCCCUAUG G CUGCCUCU	5261	AGAGGCAG GGCTAGCTACAACGA CATAGGGC	6249
2588	CUAUGGCU G CCUCUUAU	5262	CTAAGAGG GGCTAGCTACAACGA AGCCATAG	6250
2597	CCUCUUAU A CCAUGUCC	5263	GGACATGG GGCTAGCTACAACGA CTAAGAGG	6251
2600	CUUAGACC A UGUCCGGG	5264	CCCGGACA GGCTAGCTACAACGA GGTCTAAG	6252
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2612	CCGGGAAA A CCGCGGAC	5266	GTCCGCGG GGCTAGCTACAACGA TTTCCCGG	6254
2615	GGAAAACC G CGGACGCC	5267	GGCGTCCG GGCTAGCTACAACGA GGTTTTCC	6255
2619	AACCGCGG A CGCCUGGG	5268	CCCAGGCG GGCTAGCTACAACGA CCGCGGTT	6256
2621	CCGCGGAC G CCUGGGCU	5269	AGCCAGG GGCTAGCTACAACGA GTCCGCGG	6257
2627	ACGCCUGG G CUCCCAGG	5270	CCTGGGAG GGCTAGCTACAACGA CCAGGCGT	6258
2636	CUCCCAGG A CCUGCUGA	5271	TCAGCAGG GGCTAGCTACAACGA CCTGGGAG	6259
2640	CAGGACCU G CUGAACUG	5272	CAGTTCAG GGCTAGCTACAACGA AGGTCCTG	6260
2645	CCUGCUGA A CUGGUGUA	5273	TACACCAG GGCTAGCTACAACGA TCAGCAGG	6261
2649	CUGAACUG G UGUUAUGCA	5274	TGCATACA GGCTAGCTACAACGA CAGTTCAG	6262

2651	GAACUGGU G UAUGCAGA	5275	TCTGCATA GGCTAGCTACAACGA ACCAGTTC	6263
2653	ACUGGUGU A UGCAGAUU	5276	AATCTGCA GGCTAGCTACAACGA ACACCAGT	6264
2655	UGGUGUAU G CAGAUUGC	5277	GCAATCTG GGCTAGCTACAACGA ATACACCA	6265
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2662	UGCAGAUU G CCAAGGGG	5279	CCCCTTGG GGCTAGCTACAACGA AATCTGCA	6267
2671	CCAAGGGG A UGAGCUAC	5280	GTAGCTCA GGCTAGCTACAACGA CCCCTTGG	6268
2675	GGGGAUGA G CUACCUGG	5281	CCAGGTAG GGCTAGCTACAACGA TCATCCCC	6269
2678	GAUGAGCU A CCUGGAGG	5282	CCTCCAGG GGCTAGCTACAACGA AGCTCATC	6270
2687	CCUGGAGG A UGUGCGGC	5283	GCCGCACA GGCTAGCTACAACGA CCTCCAGG	6271
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2691	GAGGAUGU G CGGCUCGU	5285	ACGAGCCG GGCTAGCTACAACGA ACATCCTC	6273
2694	GAUGUGCG G CUCGUACA	5286	TGTACGAG GGCTAGCTACAACGA CGCACATC	6274
2698	UGCGGCUC G UACACAGG	5287	CCTGTGTA GGCTAGCTACAACGA GAGCCGCA	6275
2700	CGGCUCGU A CACAGGGA	5288	TCCCTGTG GGCTAGCTACAACGA ACGAGCCG	6276
2702	GCUCGUAC A CAGGGACU	5289	AGTCCCTG GGCTAGCTACAACGA GTACGAGC	6277
2708	ACACAGGG A CUUGGCCG	5290	CGGCCAAG GGCTAGCTACAACGA CCCTGTGT	6278
2713	GGGACUUG G CCGCUCGG	5291	CCGAGCGG GGCTAGCTACAACGA CAAGTCCC	6279
2716	ACUUGGCC G CUCGGAAC	5292	GTTCCGAG GGCTAGCTACAACGA GGCCAAGT	6280
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2725	CUCGGAAC G UGUGGUC	5294	GACCAGCA GGCTAGCTACAACGA GTTCCGAG	6282
2727	CGGAACGU G CUGGUCAA	5295	TTGACCAG GGCTAGCTACAACGA ACGTTCCG	6283
2731	ACGUGCUG G UCAAGAGU	5296	ACTCTTGA GGCTAGCTACAACGA CAGCACGT	6284
2738	GGUCAAGA G UCCCAACC	5297	GGTTGGGA GGCTAGCTACAACGA TCTTGACC	6285
2744	GAGUCCCA A CCAUGUCA	5298	TGACATGG GGCTAGCTACAACGA TGGGACTC	6286
2747	UCCCAACC A UGUCAAAA	5299	TTTTGACA GGCTAGCTACAACGA GGTGGA	6287
2749	CCAACCAU G UCAAAAUU	5300	AATTTTGA GGCTAGCTACAACGA ATGGTTGG	6288
2755	AUGUCAAA A UUACAGAC	5301	GTCTGTAA GGCTAGCTACAACGA TTTGACAT	6289
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2773	UCGGGCUG G CUCGGCUG	5305	CAGCCGAG GGCTAGCTACAACGA CAGCCCGA	6293
2778	CUGGCUCG G CUGCUGGA	5306	TCCAGCAG GGCTAGCTACAACGA CGAGCCAG	6294
2781	GCUCGGCU G CUGGACAU	5307	ATGTCCAG GGCTAGCTACAACGA AGCCGAGC	6295
2786	GCUCGUGG A CAUUGACG	5308	CGTCAATG GGCTAGCTACAACGA CCAGCAGC	6296
2788	UGCUGGAC A UUGACGAG	5309	CTCGTCAA GGCTAGCTACAACGA GTCCAGCA	6297
2792	GGACAUUG A CGAGACAG	5310	CTGTCTCG GGCTAGCTACAACGA CAATGTCC	6298
2797	UUGACGAG A CAGAGUAC	5311	GTA CTCTG GGCTAGCTACAACGA CTCGTCAA	6299
2802	GAGACAGA G UACCAUGC	5312	GCATGGTA GGCTAGCTACAACGA TCTGTCTC	6300
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2807	AGAGUACC A UGCAGAUG	5314	CATCTGCA GGCTAGCTACAACGA GGTA CTCT	6302
2809	AGUACCAU G CAGAUUGG	5315	CCCATCTG GGCTAGCTACAACGA ATGGTACT	6303
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2819	AGAUGGGG G CAAGGUGC	5317	GCACCTTG GGCTAGCTACAACGA CCCCATCT	6305
2824	GGGGCAAG G UGCCCAUC	5318	GATGGGCA GGCTAGCTACAACGA CTTGCC	6306
2826	GGCAAGGU G CCCAUCAA	5319	TTGATGGG GGCTAGCTACAACGA ACCTTGCC	6307
2830	AGGUGCCC A UCAAGUGG	5320	CCACTTGA GGCTAGCTACAACGA GGGCACCT	6308
2835	CCCAUCAA G UGGAUGGC	5321	GCCATCCA GGCTAGCTACAACGA TTGATGGG	6309
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2844	UGGAUGGC G CUGGAGUC	5324	GACTCCAG GGCTAGCTACAACGA GCCATCCA	6312
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2854	UGGAGUCC A UUCUCCGC	5326	GCGGAGAA GGCTAGCTACAACGA GGA CTCCA	6314

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2865	CUCCGCCG G CGGUUCAC	5328	GTGAACCG GGCTAGCTACAACGA CGGCGGAG	6316
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2876	GUUCACCC A CCAGAGUG	5331	CACTCTGG GGCTAGCTACAACGA GGGTGAAC	6319
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2900	GAGUUAUG G UGUGACUG	5338	CAGTCACA GGCTAGCTACAACGA CATAACTC	6326
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2905	AUGGUGUG A CUGUGUGG	5340	CCACACAG GGCTAGCTACAACGA CACACCAT	6328
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2923	AGCUGAUG A CUUUUGGG	5345	CCCAAAAG GGCTAGCTACAACGA CATCAGCT	6333
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2937	GGGGCCAA A CCUACGA	5347	TCGTAAGG GGCTAGCTACAACGA TTGGCCCC	6335
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2956	GGAUCCCA G CCCGGGAG	5351	CTCCCGGG GGCTAGCTACAACGA TGGGATCC	6339
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3512	GCGGUACA G UGAGGACC	5470	GGTCCTCA GGCTAGCTACAACGA TGTACCGC	6458
3518	CAGUGAGG A CCCCACAG	5471	CTGTGGGG GGCTAGCTACAACGA CCTCACTG	6459
3523	AGGACCCC A CAGUACCC	5472	GGGTACTG GGCTAGCTACAACGA GGGGTCCT	6460
3526	ACCCACA G UACCCUG	5473	CAGGGGTA GGCTAGCTACAACGA TGTGGGGT	6461
3528	CCCACAGU A CCCUGCC	5474	GGCAGGGG GGCTAGCTACAACGA ACTGTGGG	6462
3534	GUACCCCU G CCCUCUGA	5475	TCAGAGGG GGCTAGCTACAACGA AGGGGTAC	6463
3544	CCUCUGAG A CUGAUGGC	5476	GCCATCAG GGCTAGCTACAACGA CTCAGAGG	6464
3548	UGAGACUG A UGGCUACG	5477	CGTAGCCA GGCTAGCTACAACGA CAGTCTCA	6465
3551	GACUGAUG G CUACGUUG	5478	CAACGTAG GGCTAGCTACAACGA CATCAGTC	6466
3554	UGAUGGCU A CGUUGCCC	5479	GGGCAACG GGCTAGCTACAACGA AGCCATCA	6467
3556	AUGGCUAC G UUGCCCC	5480	GGGGGCAA GGCTAGCTACAACGA GTAGCCAT	6468
3559	GCUACGUU G CCCCCUG	5481	CAGGGGGG GGCTAGCTACAACGA AACGTAGC	6469
3568	CCCCCUG A CCUGCAGC	5482	GCTGCAGG GGCTAGCTACAACGA CAGGGGGG	6470

3572	CCUGACCU G CAGCCCC	5483	GGGGGCTG GGCTAGCTACAACGA AGGTCAGG	6471
3575	GACCUGCA G CCCCAGC	5484	GCTGGGGG GGCTAGCTACAACGA TGCAGGTC	6472
3582	AGCCCCCA G CCUGAAUA	5485	TATTCAGG GGCTAGCTACAACGA TGGGGGCT	6473
3588	CAGCCUGA A UAUGUGAA	5486	TTCACATA GGCTAGCTACAACGA TCAGGCTG	6474
3590	GCCUGAAU A UGUGAACC	5487	GGTTCACA GGCTAGCTACAACGA ATTCAGGC	6475
3592	CUGAAUAU G UGAACCAG	5488	CTGGTTCA GGCTAGCTACAACGA ATATTCAG	6476
3596	AUAUGUGA A CCAGCCAG	5489	CTGGCTGG GGCTAGCTACAACGA TCACATAT	6477
3600	GUGAACCA G CCAGAUGU	5490	ACATCTGG GGCTAGCTACAACGA TGGTTCAC	6478
3605	CCAGCCAG A UGUUCGGC	5491	GCCGAACA GGCTAGCTACAACGA CTGGCTGG	6479
3607	AGCCAGAU G UUCGGCCC	5492	GGGCCGAA GGCTAGCTACAACGA ATCTGGCT	6480
3612	GAUGUUCG G CCCAGCC	5493	GGCTGGGG GGCTAGCTACAACGA CGAACATC	6481
3618	CGGCCCCA G CCCCUIUC	5494	GAAGGGGG GGCTAGCTACAACGA TGGGGCCG	6482
3627	CCCCCUUC G CCCCAGAG	5495	TCTCGGGG GGCTAGCTACAACGA GAAGGGGG	6483
3638	CCGAGAGG G CCCUCUGC	5496	GCAGAGGG GGCTAGCTACAACGA CCTCTCGG	6484
3645	GGCCUCU G CCUGCUGC	5497	GCAGCAGG GGCTAGCTACAACGA AGAGGGCC	6485
3649	CUCUGCCU G CUGCCCGA	5498	TCGGGCAG GGCTAGCTACAACGA AGGCAGAG	6486
3652	UGCCUGCU G CCGACCU	5499	AGGTCGGG GGCTAGCTACAACGA AGCAGGCA	6487
3657	GCUGCCCG A CCUGCUGG	5500	CCAGCAGG GGCTAGCTACAACGA CGGGCAGC	6488
3661	CCCGACCU G CUGGUGCC	5501	GGCACCAG GGCTAGCTACAACGA AGGTCGGG	6489
3665	ACCUGCUG G UGCCACUC	5502	GAGTGGCA GGCTAGCTACAACGA CAGCAGGT	6490
3667	CUGCUGGU G CCACUCUG	5503	CAGAGTGG GGCTAGCTACAACGA ACCAGCAG	6491
3670	CUGGUGCC A CUCUGGAA	5504	TTCCAGAG GGCTAGCTACAACGA GGCACCAG	6492
3681	CUGGAAAG G CCAAGAC	5505	GTCTTGGG GGCTAGCTACAACGA CTTTCCAG	6493
3688	GGCCCAAG A CUCUCUCC	5506	GGAGAGAG GGCTAGCTACAACGA CTTGGGCC	6494
3707	AGGGAAGA A UGGGUGCG	5507	CGACCCCA GGCTAGCTACAACGA TCTTCCCT	6495
3712	AGAAUGGG G UCGUCAA	5508	TTTGACGA GGCTAGCTACAACGA CCCATTCT	6496
3715	AUGGGGUC G UCAAAGAC	5509	GTCTTTGA GGCTAGCTACAACGA GACCCCAT	6497
3722	CGUCAAAG A CGUUUUUG	5510	CAAAAACG GGCTAGCTACAACGA CTTTGACG	6498
3724	UCAAAAGAC G UUUUUGCC	5511	GGCAAAA GGCTAGCTACAACGA GTCTTTGA	6499
3730	ACGUUUUU G CCUUUGGG	5512	CCCAAAGG GGCTAGCTACAACGA AAAAACGT	6500
3740	CUUUGGGG G UGCCGUGG	5513	CCACGGCA GGCTAGCTACAACGA CCCCAAAG	6501
3742	UUGGGGGU G CCGUGGAG	5514	CTCCACGG GGCTAGCTACAACGA ACCCCCAA	6502
3745	GGGGUGCC G UGGAGAAC	5515	GTTCTCCA GGCTAGCTACAACGA GGCACCCC	6503
3752	CGUGGAGA A CCCCAGU	5516	ACTCGGGG GGCTAGCTACAACGA TCTCCACG	6504
3759	AACCCCGA G UACUUGAC	5517	GTCAAGTA GGCTAGCTACAACGA TCGGGGTT	6505
3761	CCCCGAGU A CUUGACAC	5518	GTGTCAAG GGCTAGCTACAACGA ACTCGGGG	6506
3766	AGUACUUG A CACCCAG	5519	CTGGGGTG GGCTAGCTACAACGA CAAGTACT	6507
3768	UACUUGAC A CCCAGGG	5520	CCCTGGGG GGCTAGCTACAACGA GTCAAGTA	6508
3781	AGGGAGGA G CUGCCCCU	5521	AGGGGCAG GGCTAGCTACAACGA TCCTCCCT	6509
3784	GAGGAGCU G CCCUCAG	5522	CTGAGGGG GGCTAGCTACAACGA AGCTCCTC	6510
3792	GCCCCUCA G CCCACCC	5523	GGGTGGGG GGCTAGCTACAACGA TGAGGGGC	6511
3797	UCAGCCCC A CCCUCCUC	5524	GAGGAGGG GGCTAGCTACAACGA GGGGCTGA	6512
3808	CUCCUCCU G CCUUCAGC	5525	GCTGAAGG GGCTAGCTACAACGA AGGAGGAG	6513
3815	UGCCUUCA G CCCAGCCU	5526	AGGCTGGG GGCTAGCTACAACGA TGAAGGCA	6514
3820	UCAGCCCA G CCUUCGAC	5527	GTCGAAGG GGCTAGCTACAACGA TGGGCTGA	6515
3827	AGCCUUCG A CAACCUCU	5528	AGAGGTTG GGCTAGCTACAACGA CGAAGGCT	6516
3830	CUUCGACA A CCUCUAUU	5529	AATAGAGG GGCTAGCTACAACGA TGTCGAAG	6517
3836	CAACCUCU A UUACUGGG	5530	CCCAGTAA GGCTAGCTACAACGA AGAGGTTG	6518
3839	CCUCUAUU A CUGGGACC	5531	GGTCCCAG GGCTAGCTACAACGA AATAGAGG	6519
3845	UUACUGGG A CCAGGACC	5532	GGTCCTGG GGCTAGCTACAACGA CCCAGTAA	6520
3851	GGACCAGG A CCCACCAG	5533	CTGGTGGG GGCTAGCTACAACGA CCTGGTCC	6521
3855	CAGGACCC A CCAGAGCG	5534	CGCTCTGG GGCTAGCTACAACGA GGGTCTCG	6522

3861	CCACCAGA G	CGGGGGGC	5535	GCCCCCGG	GGCTAGCTACAACGA	TCTGGTGG	6523
3868	AGCGGGGG G	CUCCACCC	5536	GGGTGGAG	GGCTAGCTACAACGA	CCCCGCT	6524
3873	GGGGCUCC A	CCCAGCAC	5537	GTGCTGGG	GGCTAGCTACAACGA	GGAGCCCC	6525
3878	UCCACCCA G	CACCUUCA	5538	TGAAGGTG	GGCTAGCTACAACGA	TGGGTGGA	6526
3880	CACCCAGC A	CCUUCAAA	5539	TTTGAAGG	GGCTAGCTACAACGA	GCTGGGTG	6527
3892	UCAAGGGG A	CACCUACG	5540	CGTAGGTG	GGCTAGCTACAACGA	CCCTTTGA	6528
3894	AAAGGGAC A	CCUACGGC	5541	GCCGTAGG	GGCTAGCTACAACGA	GTCCCTTT	6529
3898	GGACACCU A	CGGCAGAG	5542	CTCTGCCG	GGCTAGCTACAACGA	AGGTGTCC	6530
3901	CACCUACG G	CAGAGAAC	5543	GTTCTCTG	GGCTAGCTACAACGA	CGTAGGTG	6531
3908	GGCAGAGA A	CCCAGAGU	5544	ACTCTGGG	GGCTAGCTACAACGA	TCTCTGCC	6532
3915	AACCCAGA G	UACCUGGG	5545	CCCAGGTA	GGCTAGCTACAACGA	TCTGGGTT	6533
3917	CCCAGAGU A	CCUGGGUC	5546	GACCCAGG	GGCTAGCTACAACGA	ACTCTGGG	6534
3923	GUACCUGG G	UCUGGACG	5547	CGTCCAGA	GGCTAGCTACAACGA	CCAGGTAC	6535
3929	GGGUCUGG A	CGUGCCAG	5548	CTGGCACG	GGCTAGCTACAACGA	CCAGACCC	6536
3931	GUCUGGAC G	UGCCAGUG	5549	CACTGGCA	GGCTAGCTACAACGA	GTCCAGAC	6537
3933	CUGGACGU G	CCAGUGUG	5550	CACACTGG	GGCTAGCTACAACGA	ACGTCCAG	6538
3937	ACGUGCCA G	UGUGAACC	5551	GGTTCACA	GGCTAGCTACAACGA	TGGCACGT	6539
3939	GUGCCAGU G	UGAACCAG	5552	CTGGTTCA	GGCTAGCTACAACGA	ACTGGCAC	6540
3943	CAGUGUGA A	CCAGAAGG	5553	CCTTCTGG	GGCTAGCTACAACGA	TCACACTG	6541
3951	ACCAGAAG G	CCAAGUCC	5554	GGACTTGG	GGCTAGCTACAACGA	CTTCTGGT	6542
3956	AAGGCCAA G	UCCGCAGA	5555	TCTGCGGA	GGCTAGCTACAACGA	TTGGCCTT	6543
3960	CCAAGUCC G	CAGAAGCC	5556	GGCTTCTG	GGCTAGCTACAACGA	GGACTTGG	6544
3966	CCGCAGAA G	CCUGAUG	5557	CATCAGGG	GGCTAGCTACAACGA	TTCTGCGG	6545
3972	AAGCCCUG A	UGUGUCCU	5558	AGGACACA	GGCTAGCTACAACGA	CAGGGCTT	6546
3974	GCCCUGAU G	UGUCCUCA	5559	TGAGGACA	GGCTAGCTACAACGA	ATCAGGGC	6547
3976	CCUGAUGU G	UCCUCAGG	5560	CCTGAGGA	GGCTAGCTACAACGA	ACATCAGG	6548
3987	CUCAGGGA G	CAGGGAAG	5561	CTTCCCTG	GGCTAGCTACAACGA	TCCCTGAG	6549
3996	CAGGGAAG G	CCUGACUU	5562	AAGTCAGG	GGCTAGCTACAACGA	CTTCCCTG	6550
4001	AAGGCCUG A	CUUCUGCU	5563	AGCAGAAG	GGCTAGCTACAACGA	CAGGCCTT	6551
4007	UGACUUCU G	CUGGCAUC	5564	GATGCCAG	GGCTAGCTACAACGA	AGAAGTCA	6552
4011	UUCUGCUG G	CAUCAAGA	5565	TCTTGATG	GGCTAGCTACAACGA	CAGCAGAA	6553
4013	CUGCUGGC A	UCAAGAGG	5566	CCTCTTGA	GGCTAGCTACAACGA	GCCAGCAG	6554
4021	AUCAAGAG G	UGGGAGGG	5567	CCCTCCCA	GGCTAGCTACAACGA	CTCTTGAT	6555
4029	GUGGGAGG G	CCCUCCGA	5568	TCGGAGGG	GGCTAGCTACAACGA	CCTCCAC	6556
4037	GCCCUCCG A	CCACUUC	5569	GGAAGTGG	GGCTAGCTACAACGA	CGGAGGGC	6557
4040	CUCCGACC A	CUUCCAGG	5570	CCTGGAAG	GGCTAGCTACAACGA	GGTCGGAG	6558
4052	CCAGGGGA A	CCUGCCAU	5571	ATGGCAGG	GGCTAGCTACAACGA	TCCCCTGG	6559
4056	GGGAACCU G	CCAUGCCA	5572	TGGCATGG	GGCTAGCTACAACGA	AGGTTCCC	6560
4059	AACCUGCC A	UGCCAGGA	5573	TCCTGGCA	GGCTAGCTACAACGA	GGCAGGTT	6561
4061	CCUGCCAU G	CCAGGAAC	5574	GTTCTTGG	GGCTAGCTACAACGA	ATGGCAGG	6562
4068	UGCCAGGA A	CCUGUCCU	5575	AGGACAGG	GGCTAGCTACAACGA	TCCTGGCA	6563
4072	AGGAACCU G	UCCUAAGG	5576	CCTTAGGA	GGCTAGCTACAACGA	AGGTTCTT	6564
4082	CCUAAGGA A	CCUUCUU	5577	AAGGAAGG	GGCTAGCTACAACGA	TCCTTAGG	6565
4094	UCCUUCU G	CUUGAGUU	5578	AACTCAAG	GGCTAGCTACAACGA	AGGAAGGA	6566
4100	CUGCUUGA G	UUCCCAGA	5579	TCTGGGAA	GGCTAGCTACAACGA	TCAAGCAG	6567
4108	GUUCCAG A	UGGCUGGA	5580	TCCAGCCA	GGCTAGCTACAACGA	CTGGGAAC	6568
4111	CCCAGAUG G	CUGGAAGG	5581	CCTTCCAG	GGCTAGCTACAACGA	CATCTGGG	6569
4121	UGGAAGGG G	UCCAGCCU	5582	AGGCTGGA	GGCTAGCTACAACGA	CCCTTCCA	6570
4126	GGGUCCA G	CCUCGUUG	5583	CAACGAGG	GGCTAGCTACAACGA	TGGACCCC	6571
4131	CCAGCCUC G	UUGGAAGA	5584	TCTTCCAA	GGCTAGCTACAACGA	GAGGCTGG	6572
4143	GAAGAGGA A	CAGCACUG	5585	CAGTGCTG	GGCTAGCTACAACGA	TCCTCTTC	6573
4146	GAGGAACA G	CACUGGGG	5586	CCCCAGTG	GGCTAGCTACAACGA	TGTTCTTC	6574

4148	GGAACAGC A CUGGGGAG	5587	CTCCCCAG GGCTAGCTACAACGA GCTGTTCC	6575
4156	ACUGGGGA G UC UUUGUG	5588	CACAAAGA GGCTAGCTACAACGA TCCCCAGT	6576
4162	GAGUCUUU G UGGAUUCU	5589	AGAATCCA GGCTAGCTACAACGA AAAGACTC	6577
4166	CUUUGUGG A UUCUGAGG	5590	CCTCAGAA GGCTAGCTACAACGA CCACAAAG	6578
4174	AUUCUGAG G CCCUGCCC	5591	GGGCAGGG GGCTAGCTACAACGA CTCAGAAT	6579
4179	GAGGCCCU G CCCAAUGA	5592	TCATTGGG GGCTAGCTACAACGA AGGGCCTC	6580
4184	CCUGCCCA A UGAGACUC	5593	GAGTCTCA GGCTAGCTACAACGA TGGGCAGG	6581
4189	CCAAUGAG A CUCUAGGG	5594	CCCTAGAG GGCTAGCTACAACGA CTCATTGG	6582
4197	ACUCUAGG G UCCAGUGG	5595	CCACTGGA GGCTAGCTACAACGA CCTAGAGT	6583
4202	AGGGUCCA G UGGAUGCC	5596	GGCATCCA GGCTAGCTACAACGA TGGACCCT	6584
4206	UCCAGUGG A UGCCACAG	5597	CTGTGGCA GGCTAGCTACAACGA CCACTGGA	6585
4208	CAGUGGAU G CCACAGCC	5598	GGCTGTGG GGCTAGCTACAACGA ATCCACTG	6586
4211	UGGAUGCC A CAGCCCAG	5599	CTGGGCTG GGCTAGCTACAACGA GGCATCCA	6587
4214	AUGCCACA G CCCAGCUU	5600	AAGCTGGG GGCTAGCTACAACGA TGTGGCAT	6588
4219	ACAGCCCA G CUUGGCCC	5601	GGGCCAAG GGCTAGCTACAACGA TGGGCTGT	6589
4224	CCAGCUUG G CCCUUUCC	5602	GGAAAGGG GGCTAGCTACAACGA CAAGCTGG	6590
4239	CCUUCAG A UCCUGGGU	5603	ACCCAGGA GGCTAGCTACAACGA CTGGAAGG	6591
4246	GAUCCUGG G UACUGAAA	5604	TTTCAGTA GGCTAGCTACAACGA CCAGGATC	6592
4248	UCCUGGGU A CUGAAAGC	5605	GCTTTCAG GGCTAGCTACAACGA ACCCAGGA	6593
4255	UACUGAAA G CCUAGGGG	5606	CCCTAAGG GGCTAGCTACAACGA TTTCAGTA	6594
4266	UUAGGGAA G CUGGCCUG	5607	CAGGCCAG GGCTAGCTACAACGA TCCCTTAA	6595
4270	GGAAGCUG G CCUGAGAG	5608	CTCTCAGG GGCTAGCTACAACGA CAGCTTCC	6596
4284	GAGGGGAA G CGGCCCUA	5609	TAGGGCCG GGCTAGCTACAACGA TCCCTCTC	6597
4287	GGGAAGCG G CCCUAAGG	5610	CCTTAGGG GGCTAGCTACAACGA CGCTTCCC	6598
4298	CUAAGGGA G UGUCUAAG	5611	CTTAGACA GGCTAGCTACAACGA TCCCTTAG	6599
4300	AAGGGAGU G UCUAAGAA	5612	TTCTTAGA GGCTAGCTACAACGA ACTCCCTT	6600
4308	GUCUAAGA A CAAAAGCG	5613	CGCTTTTG GGCTAGCTACAACGA TCTTAGAC	6601
4314	GAACAAA G CGACCAU	5614	ATGGGTCTG GGCTAGCTACAACGA TTTGTTC	6602
4317	CAAAAGCG A CCCAUUCA	5615	TGAATGGG GGCTAGCTACAACGA CGCTTTTG	6603
4321	AGCGACCC A UUCAGAGA	5616	TCTCTGAA GGCTAGCTACAACGA GGGTCGCT	6604
4329	AUUCAGAG A CUGUCCCU	5617	AGGGACAG GGCTAGCTACAACGA CTCTGAAT	6605
4332	CAGAGACU G UCCCUGAA	5618	TTCAGGGA GGCTAGCTACAACGA AGTCTCTG	6606
4341	UCCCUGAA A CCUAGUAC	5619	GTACTAGG GGCTAGCTACAACGA TTCAGGGA	6607
4346	GAAACCUA G UACUGCCC	5620	GGGCAGTA GGCTAGCTACAACGA TAGGTTTC	6608
4348	AACCUAGU A CUGCCCCC	5621	GGGGGCAG GGCTAGCTACAACGA ACTAGGTT	6609
4351	CUAGUACU G CCCCCAU	5622	ATGGGGGG GGCTAGCTACAACGA AGTACTAG	6610
4358	UGCCCCCC A UGAGGAAG	5623	CTTCCTCA GGCTAGCTACAACGA GGGGGGCA	6611
4369	AGGAAGGA A CAGCAAUG	5624	CATTGCTG GGCTAGCTACAACGA TCCTTCCT	6612
4372	AAGGAACA G CAAUGGUG	5625	CACCATTG GGCTAGCTACAACGA TGTTCTTT	6613
4375	GAACAGCA A UGGUGUCA	5626	TGACACCA GGCTAGCTACAACGA TGCTGTTC	6614
4378	CAGCAAUG G UGUCAGUA	5627	TACTGACA GGCTAGCTACAACGA CATTGCTG	6615
4380	GCAAUGGU G UCAGUAC	5628	GATACTGA GGCTAGCTACAACGA ACCATTGC	6616
4384	UGGUGUCA G UAUCCAGG	5629	CCTGGATA GGCTAGCTACAACGA TGACACCA	6617
4386	GUGUCAGU A UCCAGGCU	5630	AGCCTGGA GGCTAGCTACAACGA ACTGACAC	6618
4392	GUAUCCAG G CUUUGUAC	5631	GTACAAAG GGCTAGCTACAACGA CTGGATAC	6619
4397	CAGGCUUU G UACAGAGU	5632	ACTCTGTA GGCTAGCTACAACGA AAAGCCTG	6620
4399	GGCUUUGU A CAGAGUGC	5633	GCACTCTG GGCTAGCTACAACGA ACAAAGCC	6621
4404	UGUACAGA G UGCUUUUC	5634	GAAAAGCA GGCTAGCTACAACGA TCTGTACA	6622
4406	UACAGAGU G CUUUUCUG	5635	CAGAAAAG GGCTAGCTACAACGA ACTCTGTA	6623
4414	GCUUUUCU G UUUAGUUU	5636	AAACTAAA GGCTAGCTACAACGA AGAAAAGC	6624
4419	UCUGUUUA G UUUUUACU	5637	AGTAAAAA GGCTAGCTACAACGA TAAACAGA	6625
4425	UAGUUUUU A CUUUUUUU	5638	AAAAAAG GGCTAGCTACAACGA AAAAACTA	6626

4434	CUUUUUUU G UUUUGUUU	5639	AAACAAAA GGCTAGCTACAACGA AAAAAAAG	6627
4439	UUUGUUUU G UUUUUUUA	5640	TAAAAAAA GGCTAGCTACAACGA AAAACAAA	6628
4451	UUUUAAAAG A UGAAAUAA	5641	TTATTTCA GGCTAGCTACAACGA CTTTAAAA	6629
4456	AAGAUGAA A UAAAGACC	5642	GGTCTTTA GGCTAGCTACAACGA TTCATCTT	6630
4462	AAAUAAAAG A CCCAGGGG	5643	CCCCTGGG GGCTAGCTACAACGA CTTTATTT	6631

Input Sequence = HSERB2R. Cut Site = R/Y
Arm Length = 8. Core Sequence = GGCTAGCTACAACGA
HSERB2R (Human c-erb-B-2 mRNA; 4473 bp)

Table V: Human HER2 Synthetic DNzyme and Target molecules

Gene	Pos	Target	Seq ID	RPI#	DNzyme	Seq ID
erbB2	377	CCACCA A UGCCAG	6632	24998	cuggca GGCTAGCTACAACGA uggugg B	6637
erbB2	766	UUCUCCG A UGUGUAA	6633	24999	uuacaca GGCTAGCTACAACGA cggagaa B	6638
erbB2	1202	UGUGCU A UGGUCU	6634	25000	agacca GGCTAGCTACAACGA agcaca B	6639
erbB2	1444	CCUCAGC G UCUUCCA	6635	25001	uggaaga GGCTAGCTACAACGA gcugagg B	6640
erbB2	1583	AUCCACC A UAACACC	6636	25002	gguguua GGCTAGCTACAACGA gguggau B	6641

A, G, C, T (*italic*) = deoxy

lower case = 2'-O-methyl

B = inverted deoxyabasic derivative

Table VI: Human HIV Hammerhead Ribozyme and Substrate Sequence

Substrate	Seq ID	Hammerhead	Seq ID
AUAAAGCU U GCCUUGAG	6642	CUCAAGGC CUGAUGAGGCCG <u>UUAGGCC</u> GAA AGCUUUUAU	6727
AGGCUAUU U UUUUAGGG	6643	CCCUAAAA CUGAUGAGGCCG <u>UUAGGCC</u> GAA AUUAGCCU	6728
GGCUAAUU U UUUAGGGA	6644	UCCCUAAA CUGAUGAGGCCG <u>UUAGGCC</u> GAA AAUAGCC	6729
GCCUCAAU A AAGCUUGC	6645	GCAAGCUU CUGAUGAGGCCG <u>UUAGGCC</u> GAA AUUGAGGC	6730
UUUCGGGU U UAUUACAG	6646	CUGUAAUA CUGAUGAGGCCG <u>UUAGGCC</u> GAA ACCCGAAA	6731
GCAGGACU C GGCUUGCU	6647	AGCAAGCC CUGAUGAGGCCG <u>UUAGGCC</u> GAA AGUCCUGC	6732

Input Sequence = HIV1. Cut Site = UH/.

Arm Length = 8. Core Sequence = CUGAUGAG GCCGUUAGGC CGAA

HIV1 Consensus

Underlined region can be any X sequence or linker, as described herein.

Table VII: Human HIV Inozyme and Substrate Sequence

Substrate	Seq ID	Inozyme	Seq ID
UGGAAAAC A GAUGGCAG	6648	CUGCCAUC CUGAUGAGGCCCGUUAGGCCGAA IUUUUCCA	6733
AAUAAAGC U UGCCUUGA	6649	UCAAGGCA CUGAUGAGGCCCGUUAGGCCGAA ICUUUAUU	6734
UCUCUAGC A GUGGCGCC	6650	GGCGCCAC CUGAUGAGGCCCGUUAGGCCGAA ICUAGAGA	6735
GGAGCCAC C CCACAAGA	6651	UCUUGUGG CUGAUGAGGCCCGUUAGGCCGAA IUGGCUCC	6736
AGUGGCGC C CGAACAGG	6652	CCUGUUCG CUGAUGAGGCCCGUUAGGCCGAA ICGCCACU	6737
GUGGCGCC C GAACAGGG	6653	CCCUGUUC CUGAUGAGGCCCGUUAGGCCGAA ICGCCAC	6738
CUCGACGC A GGACUCGG	6654	CCGAGUCC CUGAUGAGGCCCGUUAGGCCGAA ICGUCGAG	6739
CGCAGGAC U CGGCUUGC	6655	GCAAGCCG CUGAUGAGGCCCGUUAGGCCGAA IUCCUGCG	6740

Input Sequence = HIV1. Cut Site = CH/.

Arm Length = 8. Core Sequence = CUGAUGAG GCCGUUAGGC CGAA

HIV1 Consensus

Underlined region can be any X sequence or linker, as described herein.

"T" stands for Inosine.

Table VIII: Human HIV Zinzyme and Substrate Sequence

Substrate	Seq ID	Zinzyme	Seq ID
UCAAUAAA G CUUGCCUU	6656	AAGGCAAG GCCGAAAGGCGAGUGAGGUCU UUUAUUGA	6741
AGGACUCG G CUUGCUGA	6657	UCAGCAAG GCCGAAAGGCGAGUGAGGUCU CGAGUCCU	6742
GCAGUGGC G CCCGAACA	6658	UGUUCGGG GCCGAAAGGCGAGUGAGGUCU GCCACUGC	6743
CUCUAGCA G UGGCGCCC	6659	GGGCGCCA GCCGAAAGGCGAGUGAGGUCU UGCUAGAG	6744
UAGCAGUG G CGCCCGAA	6660	UUCGGGCG GCCGAAAGGCGAGUGAGGUCU CACUGCUA	6745
AGAGAUGG G UGCGAGAG	6661	CUCUCGCA GCCGAAAGGCGAGUGAGGUCU CCAUCUCU	6746
AGAUGGGU G CGAGAGCG	6662	CGCUCUCG GCCGAAAGGCGAGUGAGGUCU ACCCAUCU	6747
CUCUCGAC G CAGGACUC	6663	GAGUCCUG GCCGAAAGGCGAGUGAGGUCU GUCGAGAG	6748

Input Sequence = HIV1. Cut Site = G/Y

Arm Length = 8. Core Sequence = GCcgaagGCGaGuCaaGGuCu

HIV1 Consensus

Table IX: Human HIV DNzyme and Substrate Sequence

Substrate	Seq ID	DNzyme	Seq ID
UCAAUAAA G CUUGCCUU	6656	AAGGCAAG GGCTAGCTACAACGA TTTATTGA	6749
AGGACUCG G CUUGCUGA	6657	TCAGCAAG GGCTAGCTACAACGA CGAGTCCT	6750
GCAGUGGC G CCCGAACA	6658	TGTTCCGGG GGCTAGCTACAACGA GCCACTGC	6751
CUCUAGCA G UGGCGCCC	6659	GGGCGCCA GGCTAGCTACAACGA TGCTAGAG	6752
UAGCAGUG G CGCCCGAA	6660	TTCGGGCG GGCTAGCTACAACGA CACTGCTA	6753
AGAGAUGG G UGCGAGAG	6661	CTCTCGCA GGCTAGCTACAACGA CCATCTCT	6754
AGAUGGGU G CGAGAGCG	6662	CGCTCTCG GGCTAGCTACAACGA ACCCATCT	6755
CUCUCGAC G CAGGACUC	6663	GAGTCCTG GGCTAGCTACAACGA GTCGAGAG	6756
UAUGGAAA A CAGAUGGC	6664	GCCATCTG GGCTAGCTACAACGA TTTCCATA	6757
GAAAACAG A UGGCAGGU	6665	ACCTGCCA GGCTAGCTACAACGA CTGTTTTTC	6758
AAGCCUCA A UAAAGCUU	6666	AAGCTTTA GGCTAGCTACAACGA TGAGGCTT	6759
GGAGAGAG A UGGGUGCG	6667	CGCACCCA GGCTAGCTACAACGA CTCTCTCC	6760
GACGCAGG A CUCGGCUU	6668	AAGCCGAG GGCTAGCTACAACGA CCTGCGTC	6761

Input Sequence = HIV1. Cut Site = R/Y

Arm Length = 8. Core Sequence = GGCTAGCTACAACGA

HIV1 Consensus

Table X: Human HIV Amberzyme and Substrate Sequence

Substrate	Seq ID	Amberzyme	Seq ID
UCAAUAAA G CUUGCCUU	6656	AAGGCAAG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UUUAUUGA	6762
AGGACUCG G CUUGCUA	6657	UCAGCAAG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CGAGUCCU	6763
GCAGUGGC G CCCGAACA	6658	UGUUGGG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG GCCACUGC	6764
CUCUAGCA G UGGCGCC	6659	GGCGCCA GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UGCUAGAG	6765
UAGCAGUG G CGCCCGAA	6660	UUCGGCG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CACUGCUA	6766
AGAGAUGG G UGCGAGAG	6661	CUCUCGCA GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CCAUCUCU	6767
AGAUGGUU G CGAGAGCG	6662	CGCUCUG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG ACCCAUCU	6768
CUCUCGAC G CAGGACUC	6663	GAGUCCUG GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG GUCGAGAG	6769
GGAAACA G AUGGCAGG	6669	CCUGCCAU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UGUUUUCC	6770
AUGGGUGC G AGAGCGUC	6670	GAGGCUCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG GCAGCCAU	6771
AAAAGGG G GAUUGGG	6671	CCCCAUC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CCCCUIUU	6772
AGAAAAGG G GGAUUGG	6672	CCAAUCC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CCUUUUCU	6773
GAAGAAGG G GGAUUGG	6673	CCCAUCC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CCCUUUUC	6774
GGCUAGAA G GAGAGAGA	6674	UCUCUCUC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UUCUAGCC	6775
UUUUA AAA G AAAAGGG	6675	CCCCUUU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UUUUAAAA	6776
UAUGGCAG G AAGAAGC	6676	CGCUUCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CUGCCAU	6777
UGGCGCC G AACAGGA	6677	UCCUGUU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG GGGCGCCA	6778
GAGAGAUG G GUGCGAGA	6678	UCUGCAC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CAUCUCUC	6779
CGACGCAG G ACUGGCU	6679	AGCCGAGU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CUGCGUCG	6780
UGACUAGC G GAGGCUAG	6680	CUAGCCUC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG GCUAGUCA	6781
UAGAAGGA G AGAGAUGG	6681	CCAUCUCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UCCUUCUA	6782
AGGAGAGA G AUGGGUGC	6682	GCACCCAU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UCUCUCCU	6783
GAAGGAGA G AGAUGGU	6683	ACCCAUCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UCUCUUC	6784
UGGACGCA G GACUCGGC	6684	GCCGAGUC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG UGCGUCGA	6785
CUAGCAGU G GCGCCCGA	6685	UCGGGCGC GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG ACUGCUAG	6786
GACUAGCG G AGGCUAGA	6686	UCUAGCCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CGCUAGUC	6787
GUAGAAG G AGAGAGAU	6687	AUCUCUCU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CUUCUAGC	6788
AAAGGGG G AUUGGGG	6688	CCCCCAU GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG CCCCUIUU	6789

Input Sequence = HIV1. Cut Site = G/.

Arm Length = 8. Core Sequence = GGAGGAAACUCC CU UCAAGGACAUCGUCGCGG

HIV1 Consensus

Table XI: Human HIV Enzymatic Nucleic Acid and Target molecules

Target	Seq ID	RPI#	Enzymatic Nucleic Acid	Seq ID
GAGAUGG G UGCGAGA	6718	25003	ucucgca GGCTAGCTACAACGA ccaucuc B	6790
AUGGAAA A CAGAUGG	6719	25004	ccaucug GGCTAGCTACAACGA uuuccau B	6791
AAAACAG A UGGCAGG	6720	25005	ccugcca GGCTAGCTACAACGA cuguuuu B	6792
AGCCUCA A UAAAGCU	6721	25006	agcuuuu GGCTAGCTACAACGA ugaggcu B	6793
GAGAGAG A UGGGUGC	6722	25007	gcaccca GGCTAGCTACAACGA cucucuc B	6794
CAAUAAA G CUUGCCU	6723	25008	aggcaag gccgaaaggCgagugaGGuCu uuuaauug B	6795
GGACUCG G CUUGCUG	6724	25009	cagcaag gccgaaaggCgagugaGGuCu cgagucc B	6796
GAGAUGG G UGCGAGA	6718	25010	ucucgca gccgaaaggCgagugaGGuCu ccaucuc B	6797
GAUGGGU G CGAGAGC	6725	25011	gcucucg gccgaaaggCgagugaGGuCu acccauc B	6798
UCUCGAC G CAGGACU	6726	25012	aguccug gccgaaaggCgagugaGGuCu gucgaga B	6799

G = Guanosine

A, G, C, T (*italic*) = deoxy

lower case = 2'-O-methyl

s = phosphorothioate 3'-internucleotide linkage

C = 2'-deoxy-2'-Amino cytidine

B = inverted deoxyabasic derivative

Table XII: Human HIV-1 Sequences

Genbank Acc#	Seq Name(s)	Subtype	Organism
A04321	IIIB LAI	B	HIV-1
AF110962	96BW0402	C	HIV-1
AF110963	96BW0407	C	HIV-1
AF110968	96BW0504	C	HIV-1
AF110965	96BW0409	C	HIV-1
AF110966	96BW0410	C	HIV-1
AF110964	96BW0408	C	HIV-1
AF110975	96BW15C05	C	HIV-1
AF110974	96BW15C02	C	HIV-1
AF110973	96BW15B03	C	HIV-1
AF107771	UGSE8131	A	HIV-1
U69585	WCIPR854	B	HIV-1
U69588	WCIPR855	B	HIV-1
U69589	WCIPR9011	B	HIV-1
U69591	WCIPR9018	B	HIV-1
U69592	WCIPR9031	B	HIV-1
U69593	WCIPR9032	B	HIV-1
U69586	WCIPR8546	B	HIV-1
AF003888	NL43WC001	B	HIV-1
X01762	REHTLV3 LAI IIIB	B	HIV-1
AF075719	MNTQ MNcloneTQ	B	HIV-1
AJ239083	97CAMP645MO	MO	HIV-1
D86069	PM213	B	HIV-1
K02083	PV22	B	HIV-1
M93259	YU10	B	HIV-1
Z11530	F12CG	B	HIV-1
AB032740	TH022 95TNIH022	CRF01_AE	HIV-1
AF107770	SE7812	CRF02_AG	HIV-1
AF070521	NL43E9	B	HIV-1
AF033819	HXB2-copy LAI	B	HIV-1
AF003887	WC001	B	HIV-1
AF069140	DH123	B	HIV-1
AF110967	96BW0502	C	HIV-1
K03455	HXB2 HXB2CG	B	HIV-1
M96155	P896 89.6	B	HIV-1
X04415	MAL MALCG	ADK	HIV-1
AF133821	MB2059	D	HIV-1
D86068	MCK1	B	HIV-1
U69587	WCIPR8552	B	HIV-1
U69590	WCIPR9012	B	HIV-1
AB032741	95TNIH047 TH047	CRF01_AE	HIV-1
AB023804	93IN101	C	HIV-1
AF193275	97BL006	A	HIV-1
AF197340	90CF11697	CRF01_AE	HIV-1
AF224507	WK	B	HIV-1
AJ271445	GB8 GB8-46R	B	HIV-1
AF197338	93TH057	CRF01_AE	HIV-1
AF197339	93TH065	CRF01_AE	HIV-1
AF197341	90CF4071	CRF01_AE	HIV-1

U69584	85WCIPR54	B	HIV-1
L31963	TH475A LAI	B	HIV-1
U46016	ETH2220 C2220	C	HIV-1
U21135	WEAU160 GHOSH	B	HIV-1
AF042106	MBCC18R01	B	HIV-1
K03454	ELI	D	HIV-1
U51188	90CF402 90CR402	CRF01_AE	HIV-1
U51189	93TH253	CRF01_AE	HIV-1
U34603	H0320-2A12	B	HIV-1
M38429	JRCSF JR-CSF	B	HIV-1
M17451	RF HAT3	B	HIV-1
L02317	BC BCSG3	B	HIV-1
M93258	YU2 YU2X	B	HIV-1
M22639	Z2Z6 Z2 CDC-Z34	D	HIV-1
AF004394	AD8, AD87 ADA	B	HIV-1
AF049337	94CY032-3	CRF04_cpx	HIV-1
U34604	3202A21	B	HIV-1
L20587	ANT70	O	HIV-1
D10112	CAM1	B	HIV-1
U54771	CM240	CRF01_AE	HIV-1
U43096	D31	B	HIV-1
U37270	C18MBC	B	HIV-1
U43141	HAN	B	HIV-1
U23487	MANC	B	HIV-1
M17449	MNCG MN	B	HIV-1
L20571	MVP5180	O	HIV-1
M27323	NDK	D	HIV-1
M38431	NY5CG	B	HIV-1
M26727	OYI, 397	B	HIV-1
K02007	SF2 LAV2 ARV2	B	HIV-1
M62320	U455 U455A	A	HIV-1
U26546	WR27	B	HIV-1
AF004885	Q23	A	HIV-1
AF042100	MBC200	B	HIV-1
AF042101	MBC925	B	HIV-1
AJ006287	89SP061 89ES061	B	HIV-1
AF067154	93IN999 301999	C	HIV-1
AF067155	95IN21068 21068	C	HIV-1
AJ006022	YBF30	N	HIV-1
AF061642	SE6165 G6165	G	HIV-1
AF119820	97PVCH GR11	CRF04_cpx	HIV-1
AF119819	97PVMY GR84	CRF04_cpx	HIV-1
K02013	LAI BRU	B	HIV-1
L39106	IBNG	CRF02_AG	HIV-1
U12055	LW123	B	HIV-1
M19921	NL43 pNL43	B	HIV-1
AF061640	HH8793-1.1	G	HIV-1
AF061641	HH8793-12.1	G	HIV-1
AF063223	DJ263	CRF02_AG	HIV-1
AF049495	NC7	B	HIV-1
AF049494	499JC16	B	HIV-1
AF086817	TWCYS LM49	B	HIV-1
AF064699	BFP90	CRF06_cpx	HIV-1

AF084936	DRCBL	G	HIV-1
AF193253	VI1310 AF193253	CRF05_DF	HIV-1
AF190127	VI991	H	HIV-1
AF193276	KAL153-2	CRF03_AB	HIV-1
AF192135	BW2117	AJ	HIV-1
AJ288982	95ML127	CRF06_cpx	HIV-1
AJ288981	97SE1078	CRF06_cpx	HIV-1
AJ271370	YBF106	N	HIV-1
AJ237565	97NOGIL3	ADHK	HIV-1

CLAIMS

What we claim is:

1. A siRNA nucleic acid molecule that modulates expression of a nucleic acid molecule encoding HER2.
2. A enzymatic nucleic acid molecule that modulates expression of a nucleic acid molecule encoding HER2.
3. An enzymatic nucleic acid molecule comprising a sequence selected from the group consisting of SEQ ID NOs: 5644-6631 and 6637-6641.
4. An enzymatic nucleic acid molecule comprising at least one binding arm wherein one or more of said binding arms comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 4656-5643 and 6632-6636.
5. A siRNA nucleic acid molecule comprising a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 4656-5643 and 6632-6636.
6. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule is adapted to treat cancer.
7. The enzymatic nucleic acid molecule of any of claims 2-4, wherein said enzymatic nucleic acid molecule has an endonuclease activity to cleave RNA having HER2 sequence.
8. The enzymatic nucleic acid molecule of claim 2, wherein said enzymatic nucleic acid molecule is a DNAzyme in a 10-23 configuration.
9. The enzymatic nucleic acid molecule of claim 8, wherein said enzymatic nucleic acid molecule comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs: 4656-5643 and 6632-6636.

10. The enzymatic nucleic acid molecule of claim 8, wherein said enzymatic nucleic acid molecule comprises a sequence selected from the group consisting of SEQ ID NOs: 5644-6631 and 6637-6641.
11. The nucleic acid molecule of any of claims 1, 2, 4 or 5, wherein said nucleic acid molecule comprises between 12 and 100 bases complementary to a RNA having HER2 sequence.
12. The nucleic acid molecule of claim of any of claims 1, 2, 4 or 5, wherein said nucleic acid molecule comprises between 14 and 24 bases complementary to a RNA having HER2 sequence.
13. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule is chemically synthesized.
14. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule comprises at least one 2'-sugar modification.
15. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule comprises at least one nucleic acid base modification.
16. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule comprises at least one phosphate backbone modification.
17. A mammalian cell comprising the nucleic acid molecule of any of claims 1-5.
18. The mammalian cell of claim 17, wherein said mammalian cell is a human cell.
19. A method of reducing HER2 activity in a cell, comprising contacting said cell with the nucleic acid molecule of any of claims 1-5, under conditions suitable for said reduction of HER2 activity.
20. A method of treatment of a subject having a condition associated with the level of HER2, comprising contacting cells of said subject with the nucleic acid molecule of any of claims 1-5, under conditions suitable for said treatment.
21. The method of claim 20 further comprising the use of one or more drug therapies under conditions suitable for said treatment.

22. A method of cleaving RNA having HER2 sequence comprising contacting an enzymatic nucleic acid molecule of any of claims 2-4 with said RNA under conditions suitable for the cleavage.
23. The method of claim 22, wherein said cleavage is carried out in the presence of a divalent cation.
24. The method of claim 23, wherein said divalent cation is Mg^{2+} .
25. The nucleic acid molecule of any of claims 1-5, wherein said nucleic acid molecule comprises a cap structure, wherein the cap structure is at the 5'-end, 3'-end, or both the 5'-end and the 3'-end of said nucleic acid molecule.
26. The nucleic acid molecule of claim 25, wherein the cap structure at the 5'-end, 3'-end, or both the 5'-end and the 3'-end comprises a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative.
27. An expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of any of claims 1-5 in a manner that allows expression of the nucleic acid molecule.
28. A mammalian cell comprising an expression vector of claim 27.
29. The mammalian cell of claim 28, wherein said mammalian cell is a human cell.
30. The expression vector of claim 27, wherein said nucleic acid molecule is in a DNAzyme configuration.
31. The expression vector of claim 27, wherein said expression vector further comprises a sequence for a nucleic acid molecule complementary to a nucleic acid molecule having HER2 sequence.
32. The expression vector of claim 27, wherein said expression vector comprises a nucleic acid sequence encoding two or more of said nucleic acid molecules, which may be the same or different.
33. The expression vector of claim 32, wherein said expression vector further comprises a sequence encoding an antisense nucleic acid molecule or siRNA molecule complementary to a nucleic acid molecule having HER2 sequence.

34. A method for treatment of cancer comprising administering to a subject the nucleic acid molecule of any of claims 1-5 under conditions suitable for said treatment.
35. The method of claim 34, wherein said cancer is breast cancer.
36. The method of claim 34, wherein said cancer is ovarian cancer.
37. The method of claim 34, wherein said method further comprises administering to said subject one or more other therapies under conditions suitable for said treatment.
38. The method of claim 21 wherein said other drug therapies are chosen from monoclonal antibody therapy, chemotherapy, radiation therapy, and analgesic therapy.
39. The method of claim 37 wherein said other drug therapies are chosen from monoclonal antibody therapy, chemotherapy, radiation therapy, and analgesic therapy.
40. The method of claim 38, wherein said chemotherapy is selected from the group consisting of paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, and vinorelbine.
41. The method of claim 38, wherein said monoclonal antibody is Herceptin (trastuzumab).
42. The method of claim 39, wherein said chemotherapy is selected from the group consisting of paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, and vinorelbine.
43. The method of claim 39, wherein said monoclonal antibody is Herceptin (trastuzumab).
44. A composition comprising a nucleic acid molecule of any of claims 1-5 in a pharmaceutically acceptable carrier.

45. A method of administering to a cell a nucleic acid molecule of any of claims 1-5 comprising contacting said cell with the nucleic acid molecule under conditions suitable for said administration.
46. The method of claim 45, wherein said cell is a mammalian cell.
47. The method of claim 45, wherein said cell is a human cell.
48. The method of claim 45, wherein said administration is in the presence of a delivery reagent.
49. The method of claim 48, wherein said delivery reagent is a lipid.
50. The method of claim 49, wherein said lipid is a cationic lipid.
51. The method of claim 49, wherein said lipid is a phospholipid.
52. The method of claim 48, wherein said delivery reagent is a liposome.
53. A siRNA nucleic acid molecule that modulates expression of a nucleic acid molecule encoding K-Ras.
54. A siRNA nucleic acid molecule that modulates expression of a nucleic acid molecule encoding H-Ras or N-Ras.
55. An enzymatic nucleic acid molecule that modulates expression of a nucleic acid molecule encoding K-Ras.
56. An enzymatic nucleic acid molecule that modulates expression of a nucleic acid molecule encoding H-Ras or N-Ras.
57. An enzymatic nucleic acid molecule comprising a sequence of SEQ ID NOs: 2329-4655.
58. An enzymatic nucleic acid molecule comprising at least one binding arm wherein one or more of said binding arms comprises a sequence complementary to a sequence of SEQ ID NOs: 1-2328.
59. A siRNA nucleic acid molecule comprising a sequence complementary to a sequence of SEQ ID NOs: 1-2328.

60. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule is adapted to treat cancer.
61. The enzymatic nucleic acid molecule of any of claims 55, 57 or 58, wherein said enzymatic nucleic acid molecule has an endonuclease activity to cleave RNA having a K-Ras sequence.
62. The enzymatic nucleic acid molecule of any of claims 56-58, wherein said enzymatic nucleic acid molecule has an endonuclease activity to cleave RNA having an H-Ras sequence.
63. The enzymatic nucleic acid molecule of claim 55 or claim 56, wherein said enzymatic nucleic acid molecule is a DNAzyme in a 10-23 configuration.
64. The enzymatic nucleic acid molecule of claim 63, wherein said enzymatic nucleic acid molecule comprises a sequence complementary to a sequence of SEQ ID NOs: 1-2328.
65. The enzymatic nucleic acid molecule of claim 63, wherein said enzymatic nucleic acid molecule comprises a sequence of SEQ ID NOs: 2329-4655.
66. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule comprises between 12 and 100 bases complementary to an RNA having K-Ras, H-Ras and/or N-Ras sequence.
67. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule comprises between 14 and 24 bases complementary to an RNA having K-Ras, H-Ras and/or N-Ras sequence.
68. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule is chemically synthesized.
69. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule comprises at least one 2'-sugar modification.
70. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule comprises at least one nucleic acid base modification.
71. The nucleic acid molecule of any of claims 53-59, wherein said enzymatic nucleic acid molecule comprises at least one phosphate backbone modification.

72. A mammalian cell comprising the nucleic acid molecule of any of claims 53-59.
73. The mammalian cell of claim 72, wherein said mammalian cell is a human cell.
74. A method of reducing K-Ras activity in a cell, comprising contacting said cell with the nucleic acid molecule of any of claims 53, 55, 57, 58 or 59, under conditions suitable for said reduction of K-Ras activity.
75. A method of reducing H-Ras activity in a cell, comprising contacting said cell with the nucleic acid molecule of any of claims 54, 56, 57, 58 or 59, under conditions suitable for said reduction of H-Ras activity.
76. A method of treatment of a subject having a condition associated with the level of K-Ras, comprising contacting cells of said subject with the nucleic acid molecule of any of claims 53, 55, 57, 58 or 59, under conditions suitable for said treatment.
77. A method of treatment of a subject having a condition associated with the level of H-Ras, comprising contacting cells of said subject with the nucleic acid molecule of any of claims 54, 56, 57, 58 or 59, under conditions suitable for said treatment.
78. The method of claim 76 further comprising the use of one or more drug therapies under conditions suitable for said treatment.
79. The method of claim 77 further comprising the use of one or more drug therapies under conditions suitable for said treatment.
80. A method of cleaving RNA having a K-Ras sequence comprising contacting an nucleic acid molecule of any of claims 53, 55, 57, 58 or 59, with said RNA under conditions suitable for the cleavage.
81. A method of cleaving RNA having a H-Ras sequence comprising contacting an nucleic acid molecule of any of claims 54, 56, 57, 58 or 59, with said RNA under conditions suitable for the cleavage.
82. The method of claim 80, wherein said cleavage is carried out in the presence of a divalent cation.

83. The method of claim 81, wherein said cleavage is carried out in the presence of a divalent cation.
84. The method of claim 82, wherein said divalent cation is Mg^{2+} .
85. The method of claim 83, wherein said divalent cation is Mg^{2+} .
86. The nucleic acid molecule of any of claims 53-59, wherein said nucleic acid molecule comprises a cap structure, wherein the cap structure is at the 5'-end, 3'-end, or both the 5'-end and the 3'-end of said nucleic acid molecule.
87. The nucleic acid molecule of claim 86, wherein the cap structure comprises a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative.
88. An expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of any of claims 53-59 in a manner that allows expression of the nucleic acid molecule.
89. A mammalian cell comprising an expression vector of claim 88.
90. The mammalian cell of claim 89, wherein said mammalian cell is a human cell.
91. The expression vector of claim 88, wherein said nucleic acid molecule is in a DNAzyme configuration.
92. The expression vector of claim 88, wherein said expression vector further comprises a sequence for a nucleic acid molecule complementary to a nucleic acid molecule having a K-Ras sequence.
93. The expression vector of claim 88, wherein said expression vector further comprises a sequence for a nucleic acid molecule complementary to a nucleic acid molecule having a H-Ras sequence.
94. The expression vector of claim 88, wherein said expression vector comprises a nucleic acid sequence encoding two or more of said nucleic acid molecules, which may be the same or different.
95. The expression vector of claim 88, wherein said expression vector further comprises a sequence encoding an antisense nucleic acid molecule or siRNA

nucleic acid molecule complementary to a nucleic acid molecule having a K-Ras sequence.

96. The expression vector of claim 88, wherein said expression vector further comprises a sequence encoding an antisense nucleic acid molecule or siRNA nucleic acid molecule complementary to a nucleic acid molecule having a H-Ras sequence.
97. A method for the treatment of cancer comprising administering to a subject the nucleic acid molecule of any of claims 53-59 under conditions suitable for said treatment.
98. The method of claim 97, wherein said cancer is colorectal cancer.
99. The method of claim 97, wherein said cancer is lung cancer.
100. The method of claim 97, wherein said cancer is prostate cancer.
101. The method of claim 97, wherein said cancer is bladder cancer.
102. The method of claim 97, wherein said cancer is breast cancer.
103. The method of claim 97, wherein said cancer is pancreatic cancer.
104. The method of claim 97, wherein said method further comprises administering to said patient one or more other therapies under conditions suitable for said treatment.
105. The method of claim 78 wherein said other drug therapies are chosen from monoclonal antibody therapy, chemotherapy, radiation therapy, and analgesic therapy.
106. The method of claim 79 wherein said other drug therapies are chosen from monoclonal antibody therapy, chemotherapy, radiation therapy, and analgesic therapy.
107. The method of claim 104 wherein said other drug therapies are chosen from monoclonal antibody therapy, chemotherapy, radiation therapy, and analgesic therapy.

108. The method of claim 105, wherein said chemotherapy is selected from the group consisting of paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, and vinorelbine.
109. The method of claim 106, wherein said chemotherapy is selected from the group consisting of paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, and vinorelbine.
110. The method of claim 107, wherein said chemotherapy is selected from the group consisting of paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubin, fluorouracil carboplatin, edatrexate, gemcitabine, and vinorelbine.
111. A composition comprising a nucleic acid molecule of any of claims 53-59 and a pharmaceutically acceptable carrier.
112. A method of administering to a cell a nucleic acid molecule of any of claims 53-59 comprising contacting said cell with the enzymatic nucleic acid molecule under conditions suitable for said administration.
113. The method of claim 112, wherein said cell is a mammalian cell.
114. The method of claim 113, wherein said cell is a human cell.
115. The method of claim 112, wherein said administration is in the presence of a delivery reagent.
116. The method of claim 115, wherein said delivery reagent is a lipid.
117. The method of claim 116, wherein said lipid is a cationic lipid.
118. The method of claim 116, wherein said lipid is a phospholipid.
119. The method of claim 115, wherein said delivery reagent is a liposome.
120. A siRNA nucleic acid molecule which modulates expression of a nucleic acid molecule encoding HIV or a component of HIV.

121. An enzymatic nucleic acid molecule which modulates expression of a nucleic acid molecule encoding HIV or a component of HIV, wherein said enzymatic nucleic acid molecule is in an Inozyme, G-cleaver, Zinzyme or Amberzyme configuration.
122. An enzymatic nucleic acid molecule comprising a sequence selected from the group consisting of SEQ ID NOs. 6727-6799.
123. An enzymatic nucleic acid molecule comprising at least one binding arm wherein one or more of said binding arms comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6642-6726.
124. A siRNA nucleic acid molecule comprising a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6642-6726.
125. The nucleic acid of any of claims 120-124, wherein said nucleic acid molecule is adapted to HIV infection or acquired immunodeficiency syndrome (AIDS).
126. The enzymatic nucleic acid molecule of any of claims 121-123, wherein said enzymatic nucleic acid molecule has an endonuclease activity to cleave RNA having a HIV sequence.
127. The enzymatic nucleic acid molecule of claim 121, wherein said enzymatic nucleic acid molecule is in an Inozyme configuration.
128. The enzymatic nucleic acid molecule of claim 121, wherein said enzymatic nucleic acid molecule is in a Zinzyme configuration.
129. The enzymatic nucleic acid molecule of claim 121, wherein said enzymatic nucleic acid molecule is in a G-cleaver configuration.
130. The enzymatic nucleic acid molecule of claim 121, wherein said enzymatic nucleic acid molecule is in an Amberzyme configuration.
131. The enzymatic nucleic acid molecule of claim 123, wherein said enzymatic nucleic acid molecule is in a DNAzyme configuration.
132. The enzymatic nucleic acid molecule of claim 123, wherein said enzymatic nucleic acid molecule is in a Hammerhead configuration.

133. The enzymatic nucleic acid molecule of claim 127, wherein said Inozyme comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6648-6655.
134. The enzymatic nucleic acid molecule of claim 127, wherein said Inozyme comprises a sequence selected from the group consisting of SEQ ID NOs. 6733-6740.
135. The enzymatic nucleic acid molecule of claim 128, wherein said Zinzyme comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6663 and 6723-6726.
136. The enzymatic nucleic acid molecule of claim 128, wherein said Zinzyme comprises a sequence selected from the group consisting of SEQ ID NOs. 6741-6748 and 6795-6799.
137. The enzymatic nucleic acid molecule of claim 130, wherein said Amberzyme comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6688.
138. The enzymatic nucleic acid molecule of claim 130, wherein said Amberzyme comprises a sequence selected from the group consisting of SEQ ID NOs. 6762-6789.
139. The enzymatic nucleic acid molecule of claim 131, wherein said DNAzyme comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6656-6668 and 6718-6722.
140. The enzymatic nucleic acid molecule of claim 131, wherein said DNAzyme comprises a sequence selected from the group consisting of SEQ ID NOs. 6749-6761 and 6790-6794.
141. The enzymatic nucleic acid molecule of claim 132, wherein said Hammerhead comprises a sequence complementary to a sequence selected from the group consisting of SEQ ID NOs. 6642-6647.
142. The enzymatic nucleic acid molecule of claim 132, wherein said Hammerhead comprises a sequence selected from the group consisting of SEQ ID NOs 6727-6732.

143. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises between 12 and 100 bases complementary to a nucleic acid molecule encoding HIV.
144. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises between 14 and 24 bases complementary to a nucleic acid molecule encoding HIV.
145. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule is chemically synthesized.
146. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises at least one 2'-sugar modification.
147. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises at least one nucleic acid base modification.
148. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises at least one phosphate backbone modification.
149. A mammalian cell comprising the nucleic acid molecule of any of claims 120-124
150. The mammalian cell of claim 149, wherein said mammalian cell is a human cell.
151. A method of reducing HIV activity in a cell, comprising contacting said cell with the nucleic acid molecule of any of claims 120-124, under conditions suitable for said reduction of HIV activity.
152. A method of treatment of a subject having a condition associated with the level of HIV, comprising contacting cells of said subject with the nucleic acid molecule of any of claims 120-124, under conditions suitable for said treatment.
153. The method of claim 151 further comprising the use of one or more drug therapies under conditions suitable for said treatment.
154. The method of claim 152 further comprising the use of one or more drug therapies under conditions suitable for said treatment.

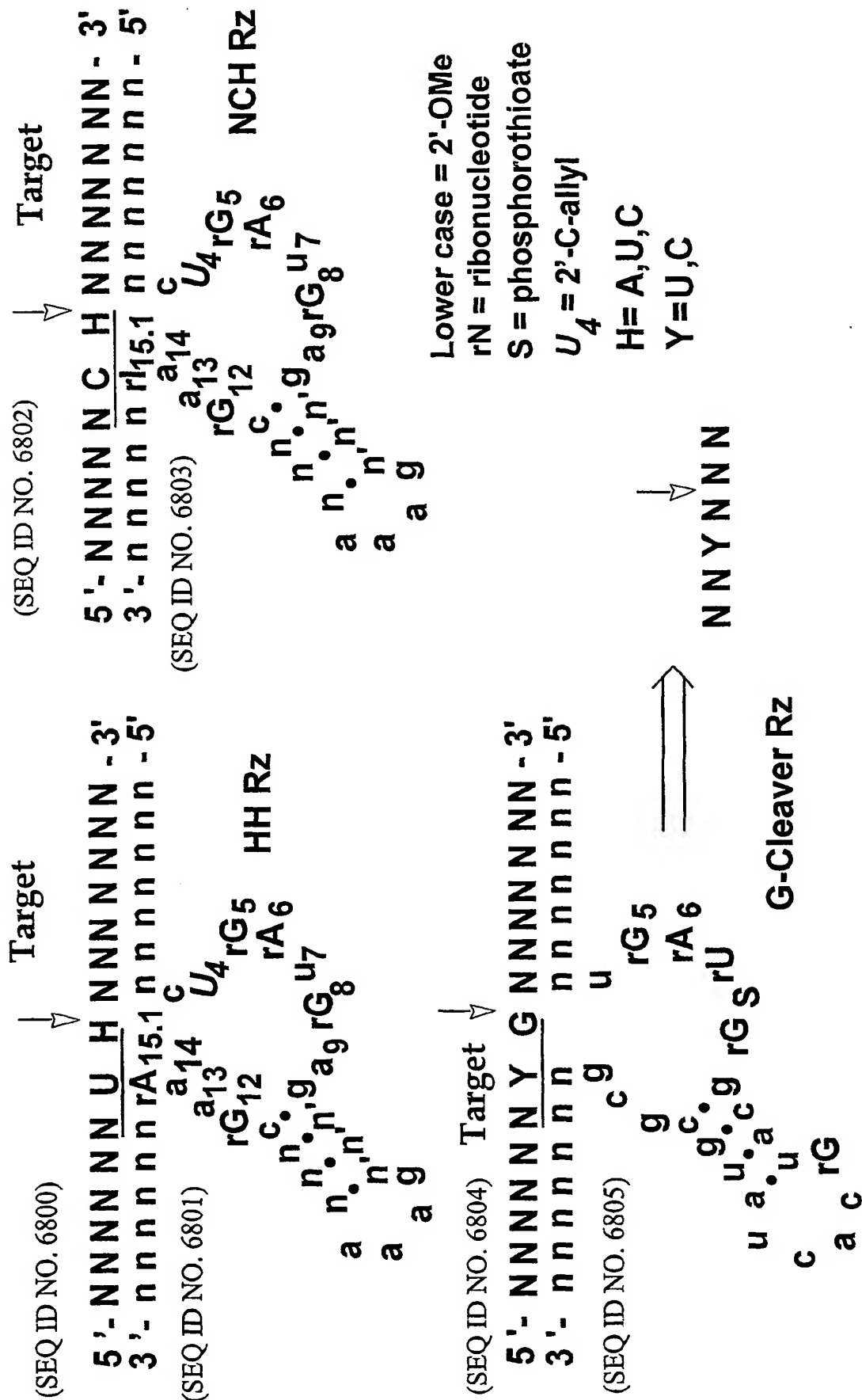
155. A method of cleaving RNA of an HIV gene comprising contacting an enzymatic nucleic acid molecule of any of claims 121-123 with said RNA of a HIV gene under conditions suitable for the cleavage.
156. The method of claim 155, wherein said cleavage is carried out in the presence of a divalent cation.
157. The method of claim 156, wherein said divalent cation is Mg^{2+} .
158. The nucleic acid molecule of any of claims 120-124, wherein said nucleic acid molecule comprises a cap structure, wherein the cap structure is at the 5'-end, 3'-end, or both the 5'-end and the 3'-end of said nucleic acid molecule.
159. The nucleic acid molecule of claim 158, wherein the cap structure at the 5'-end, 3'-end, or both the 5'-end and the 3'-end comprises a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative.
160. An expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of any of claims 120-124 in a manner which allows expression of the nucleic acid molecule.
161. A mammalian cell comprising an expression vector of claim 160.
162. The mammalian cell of claim 161, wherein said mammalian cell is a human cell.
163. An expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of any of claims 122 or 123 in a manner which allows expression of the nucleic acid molecule, wherein said nucleic acid molecule is in a hammerhead configuration.
164. The expression vector of claim 160, wherein said expression vector further comprises a sequence for a nucleic acid molecule complementary to the RNA of HIV.
165. The expression vector of claim 160, wherein said expression vector comprises a nucleic acid sequence encoding two or more of said nucleic acid molecules, which may be the same or different.

166. The expression vector of claim 165, wherein said expression vector further comprises a sequence encoding a siRNA nucleic acid molecule complementary to the RNA of HIV gene.
167. A method for treatment of acquired immunodeficiency syndrome (AIDS) or an AIDS related condition comprising administering to a subject the nucleic acid molecule of any of claims 120-124 under conditions suitable for said treatment.
168. The method of claim 167, wherein said AIDS related condition is Kaposi's sarcoma, lymphoma, cervical cancer, squamous cell carcinoma, cardiac myopathy, rheumatic disease, or opportunistic infection.
169. The method of claim 167, wherein said method further comprises administering to said subject one or more other therapies.
170. The nucleic acid molecule of claim 121 or claim 123, wherein said nucleic acid molecule comprises at least five ribose residues, at least ten 2'-O-methyl modifications, and a 3'- end modification.
171. The nucleic acid molecule of claim 170, wherein said nucleic acid molecule further comprises phosphorothioate linkages on at least three of the 5' terminal nucleotides.
172. The nucleic acid molecule of claim 170, wherein said 3'- end modification is a 3'-3' inverted abasic moiety.
173. The method of claim 153 wherein said other drug therapies chosen from antiviral therapy, monoclonal antibody therapy, chemotherapy, radiation therapy, analgesic therapy, and anti-inflammatory therapy.
174. The method of claim 173, wherein said antiviral therapy is chosen from treatment with AZT, ddC, ddI, d4T, 3TC, Ribavirin, delvaridine, nevirapine, efavirenz, ritonavir, saquinivir, indinavir, amprenivir, nelfinavir, and lopinavir.
175. The method of claim 154 wherein said other drug therapies are chosen from antiviral therapy, monoclonal antibody therapy, chemotherapy, radiation therapy, analgesic therapy, and anti-inflammatory therapy.

176. The method of claim 175, wherein said antiviral therapy is chosen from treatment with AZT, ddC, ddI, d4T, 3TC, Ribavirin, delvaridine, nevirapine, efavirenz, ritonavir, saquinivir, indinavir, amprenivir, nelfinavir, and lopinavir.
177. The method of claim 169 wherein said other drug therapies are chosen from antiviral therapy, monoclonal antibody therapy, chemotherapy, radiation therapy, analgesic therapy, and anti-inflammatory therapy.
178. The method of claim 177, wherein said antiviral therapy is chosen from treatment with AZT, ddC, ddI, d4T, 3TC, Ribavirin, delvaridine, nevirapine, efavirenz, ritonavir, saquinivir, indinavir, amprenivir, nelfinavir, and lopinavir.
179. A pharmaceutical composition comprising a nucleic acid molecule of any of claims 120-124 in a pharmaceutically acceptable carrier.
180. The nucleic acid molecule of claim 120 or 121, wherein said component of HIV is nef.
181. The nucleic acid molecule of claim 120 or 121, wherein said component of HIV is vif.
182. The nucleic acid molecule of claim 120 or 121, wherein said component of HIV is tat.
183. The nucleic acid molecule of claim 120 or 121, wherein said component of HIV is rev.
184. The nucleic acid molecule of claim 120 or 121, wherein said component of HIV is LTR.
185. The nucleic acid molecule of claim 184, wherein said LTR is the 3'-LTR.
186. The nucleic acid molecule of claim 184, wherein said LTR is the 5'-LTR.
187. A method of administering to a cell a nucleic acid molecule of any of claims 120-124 comprising contacting said cell with the nucleic acid molecule under conditions suitable for said administration.
188. The method of claim 187, wherein said cell is a mammalian cell.

189. The method of claim 187, wherein said cell is a human cell.
190. The method of claim 187, wherein said administration is in the presence of a delivery reagent.
191. The method of claim 190, wherein said delivery reagent is a lipid.
192. The method of claim 191, wherein said lipid is a cationic lipid.
193. The method of claim 191, wherein said lipid is a phospholipid.
194. The method of claim 190, wherein said delivery reagent is a liposome.

Figure 1: Examples of Nuclease Stable Ribozyme Motifs



**Figure 2: 2'-O-Me substituted Amberzyme
Enzymatic Nucleic Acid Motif**

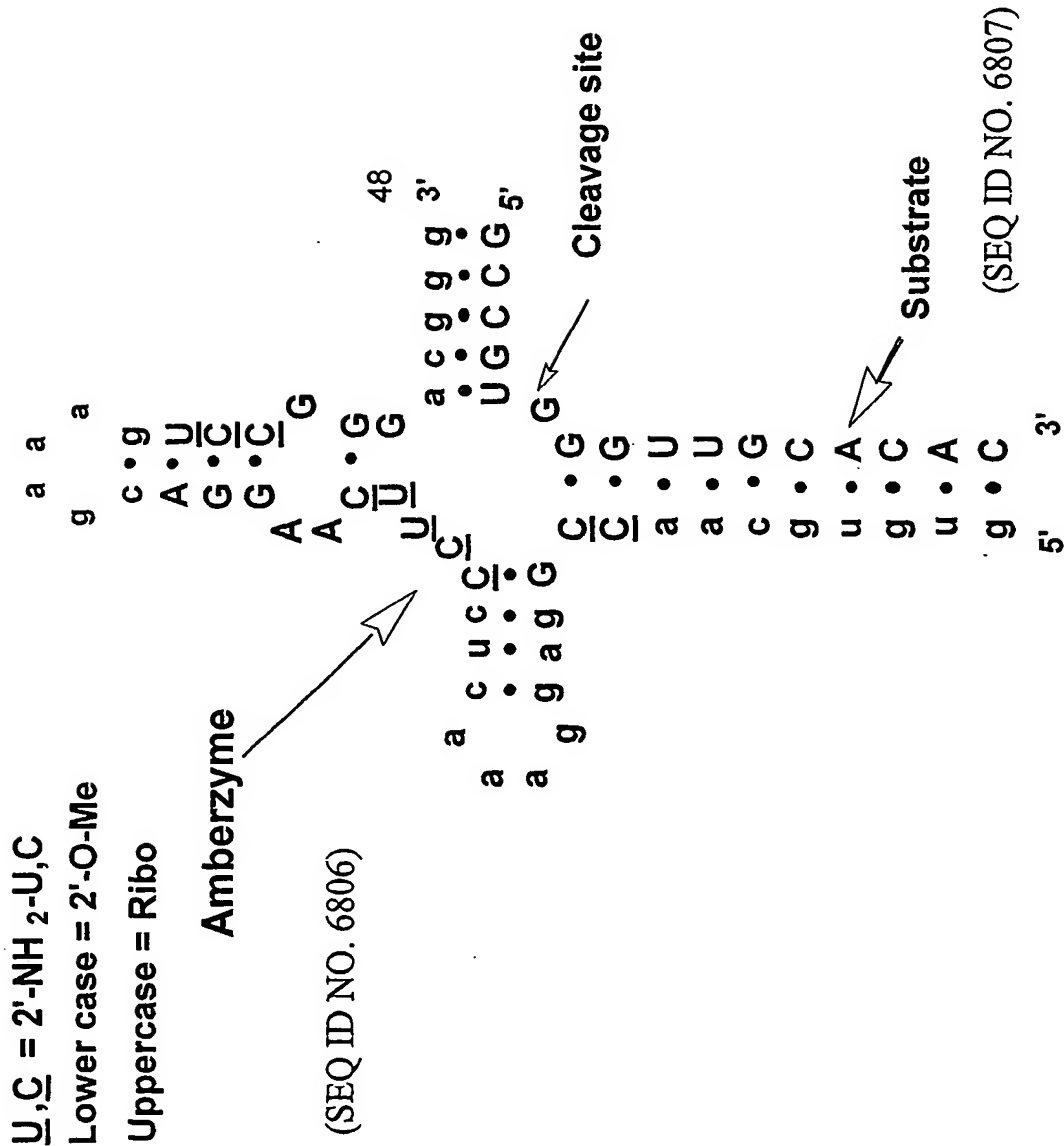
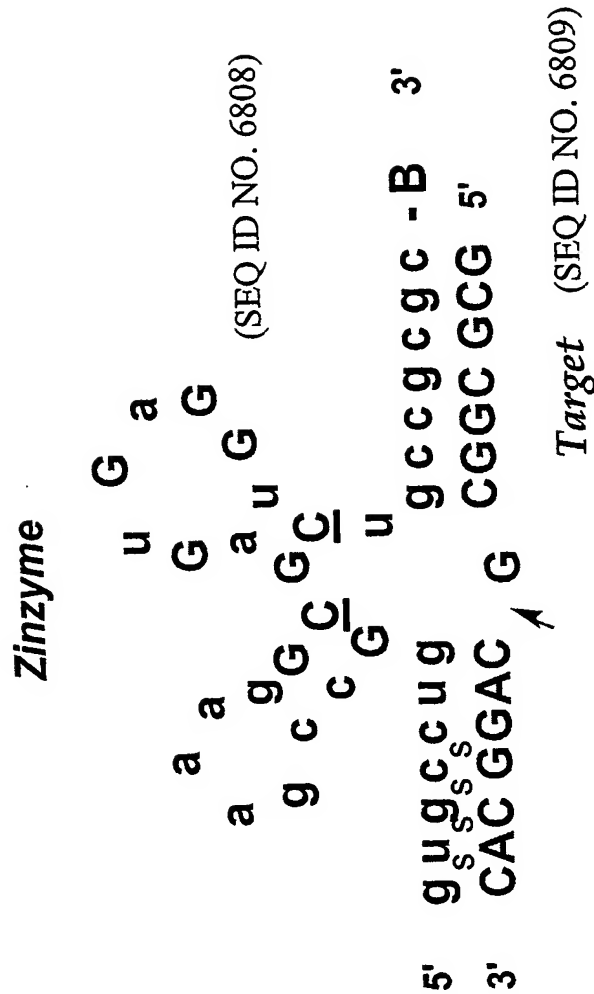


Figure 3: Stabilized Zinzyme Ribozyme Motif



Legend

Uppercase: indicates natural ribo residues

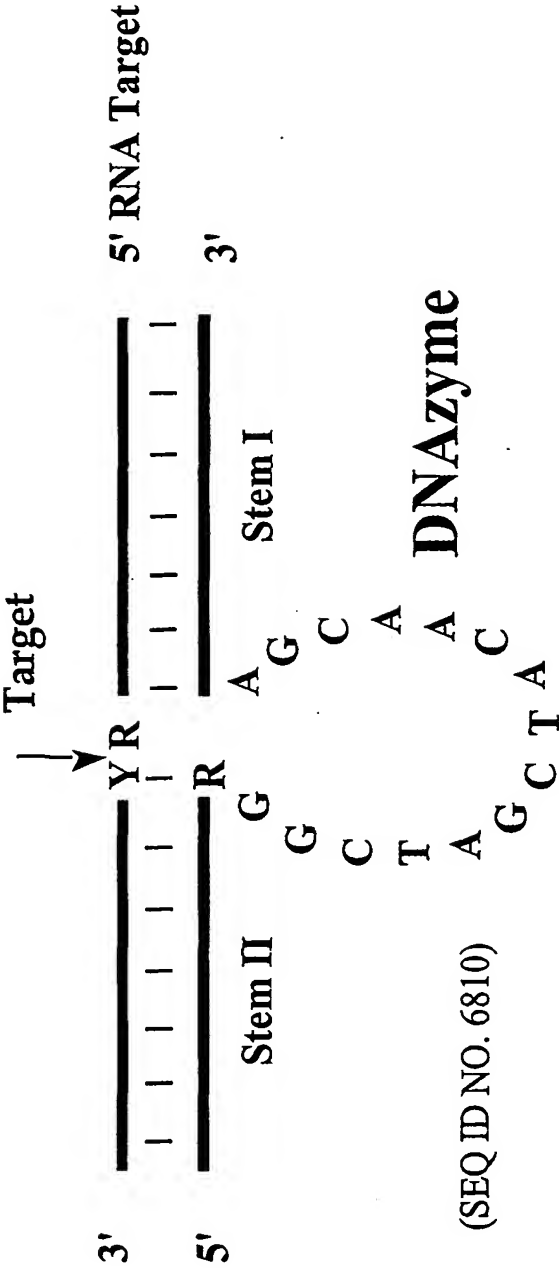
C : indicates 2'-deoxy-2'-amino Cytidine

Lowercase: 2'-O-methyl

S: phosphorothioate/phosphorodithioate linkage

B: 3'-3' abasic moiety

Figure 4: DNzyme Motif



Legend
Y = U or C
R = A or G

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(54) Title: NUCLEIC ACID TREATMENT OF DISEASES OR CONDITIONS RELATED TO LEVELS OF RAS, HER2 AND HIV

(57) Abstract: The present invention relates to nucleic acid molecules, including enzymatic nucleic acid molecules, such as DNazymes (e.g. DNA enzymes, catalytic DNA), siRNA, aptamers, and antisense that modulate the expression of Ras genes such as K-Ras, H-Ras, and/or N-Ras, HIV genes such as HIV-1, and HER2 genes.

WO 02/097114 A3

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/16840

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) : C12N 5/00, C07H 21/04

US CL : 435/366; 435/363, 536/23.2, 536/24.5

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/366; 435/363, 536/23.2, 536/24.5

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
Biosis, Medline, Scisearch, CA, and USPTO sequence search databases.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 99/31118 A1 (GEORGETOWN UNIVERSITY) 24 June 1999 (24.06.1999),	2, 6, 7, 11-13, 19-37,
—	Abstract, specification throughout.	and 44-52
Y		8, 38-43
Y	US 5,968,748 A (BENNETT et al) 19 October 1999 (19.10.1999), throughout.	1, 6, 11-52
Y	US 5,910,583 A (MARKS et al) 08 June 1999 (08.06.1999). Throughout.	1, 6, 11-52
Y	NISHIKURA, K. A Short Primer on RNAi: RNA-directed RNA Polymerase Acts as a	1, 6, 11-52
—	Key Catalyst. Cell. 16 November 2001, Vol. 107, pages 415-418, throughout.	1, 6, 11-52
A		
Y	COUSENS, L. et al. Tyrosine Kinase Receptor with Extensive Homology to EGF	1, 6,
—	Receptor Shares Chromosomal Location with neu Oncogene. Science. December 1985,	
A	Vol. 230, pages 1132-1139. Particularly figure 1.	1, 6,

☐ Further documents are listed in the continuation of Box C.

☐ See patent family annex.

* Special categories of cited documents:

"A" document defining the general state of the art which is not considered to be of particular relevance

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"&" document member of the same patent family

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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/16840

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)

This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claim Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:
2. ☐ Claim Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. ☐ Claim Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:
Please See Continuation Sheet

1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
 2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
 3. ☒ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-4,6-8 and 11-52
 4. ☐ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
- Remark on Protest ☐ The additional search fees were accompanied by the applicant's protest.
☒ No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

PCT/US02/16840

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING:

Group 1-988, 989-993, 4966-7292, and 14276-14348 drawn to a ribozyme comprising SEQ ID NOS. 5644-6631, 6637-6641, 2329-4655, and 6727-6799 respectively, and claims respectively drawn to these.

Groups 994-1981, 1982-1986, 7293-9620, and 14349-14433, drawn to ribozymes targeted to a sequence comprising any one of SEQ ID NOS. 4656-5643, 6632-6636, 1-2328, and 6642-6726 respectively, and claims respectively drawn to these.

Groups 1987-2974, 2975-2979, and 9621-11948, 14434-14518, drawn to siRNA targeted to a sequence comprising any one of SEQ ID NOS. 4656-5643, 6632-6636, 1-2328 and 6642-6726 respectively, and claims respectively drawn to these.

Groups 2980-3967, 3968-3972, and 11949-14275, drawn to a DNAzyme targeted to a sequence comprising any one of SEQ ID NOS. 4656-5643, 6632-6636, and 2329-4655 respectively, and claims respectively drawn to these.

Groups 3973-4960, and 4961-4965, drawn to a DNAzyme comprising SEQ ID NOS. 5644-6631, and 6637-6641 respectively, and claims respectively drawn to these.

Groups 14519-14526, drawn to the inozyme targeted to the sequence comprising any one of SEQ ID NOS 6727-6799, respectively.

Groups 14527-14534, drawn to the inozyme of SEQ ID NOS 6733-6740, respectively.

Groups 14535-14542 and 14543-14546, drawn to the zinzyme targeted to the sequence comprising any one of SEQ ID NOS 6656-6663, and 6723-6726, respectively.

Groups 14547-14554, and 14555-14559, drawn to the zinzyme of SEQ ID NOS 6741-6748, and 6795-6799, respectively.

Groups 14560-14592, drawn to the amberzyme targeted to the sequence comprising any one of SEQ ID NOS 6656-6688, respectively.

Groups 14593-14620, drawn to the amberzyme of SEQ ID NOS 6762-6789, respectively.

Groups 14621-14633, and 14634-14638, drawn to the DNAzyme targeted to the sequence comprising any one of SEQ ID NOS 6749-6761 and 6790-6794, respectively.

Groups 14639-14651, and 14652-14656, drawn to the DNAzyme of SEQ ID NOS 6749-6761, and 6790-6794, respectively.

Groups 14657-14662, drawn to the hammerhead ribozyme targeted to the sequence comprising any one of SEQ ID NOS 6642-6647, respectively.

Groups 14663-14668, drawn to the hammerhead ribozyme of SEQ ID NOS 6727-6732, respectively.

This International Searching Authority considers that the international application does not comply with the requirements of unity of invention (Rules 13.1, 13.2 and 13.3) for the reasons indicated below:

The claims of the instant inventions are drawn to siRNA and enzymatic nucleic acids comprising ribozymes and DNAzymes, and including the motif-types hammerhead, hairpin, hepatitis Delta virus, group I intron, VS nucleic acid, amberzyme, zinzyme, RNase P nucleic acid, NCH motif, and G-cleaver. All are directed to targets comprising Ras isoforms, the HER2 protein, and any protein encoded by the HIV virus. These targets and effector molecules comprise several thousand nucleotide sequences.

This international searching authority considers that the international application does not comply with the requirements of unity of invention (Rules 13.1, 13.2, and 13.3) for the reasons indicated below:

INTERNATIONAL SEARCH REPORT

According to the guidelines in Section (f)(i)(a) of Annex B of the PCT Administrative Instructions, the special technical feature as defined by PCT Rule 13.2 shall be considered to be met when all the alternatives of a Markush-group are of similar nature. For chemical alternatives, such as the claimed sequences, the Markush group shall be regarded as being of similar nature when

(A) all alternatives have a common property or activity and

(B)(1) a common structure is present, i.e., a significant structure is shared by all of the alternatives or

(B)(2) in cases where the common structure cannot be the unifying criteria, all alternatives belong to an art recognized class of compounds in the art to which the invention pertains.

The instant sequences are considered to be each separate inventions for the following reasons:

The sequences do not meet the criteria of (A), common property or activity and (B)(1), they do not share, one with another, a common core structure. Accordingly, unity of invention between the antisense sequences is lacking and each sequence claimed is considered to constitute a special technical feature. The sequences each behave in a different way in the context of the claimed invention. Each member of the class cannot be substituted, one for the other, with the expectation that the same intended result would be achieved.

Although the ribozyme or antisense sequences or targets listed above each target and modulate expression of any Ras isoforms, the HER2 protein, and any nucleotide encoding a protein of the HIV virus, each ribozyme sequence and/or motif, and each antisense sequence is considered to be unrelated, since each ribozyme or antisense oligo claimed is structurally and functionally independent and distinct for the following reasons: each ribozyme or antisense oligo has a unique nucleotide sequence, each ribozyme or antisense oligo targets a different and specific region of their respective genes, and each ribozyme or antisense oligo, upon binding to the transcript, functionally modulates (increases or decreases) the expression of the gene. Furthermore, a search of more than one (1) of the ribozyme or antisense oligo sequences in this application presents an undue burden on the Patent and Trademark Office due to the complex nature of the search and corresponding examination of more than one (1) of the claimed ribozyme or antisense oligo sequences. In view of the foregoing, one (1) target sequence from the above listed claims and its corresponding antisense OR ribozyme sequence (regardless of motif-type) is considered to be a reasonable number of sequences for examination. Accordingly, applicants are required to elect one (1) target sequence and its corresponding siRNA OR ribozyme sequence from claims to be searched in the instant application.

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